



# Effects of Renal Disease on Pharmacokinetics

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# **GOALS of Effects of Renal Disease on Pharmacokinetics Lecture**

**A. Dose Adjustment in patients with renal Impairment**

**B. Effect of Renal Disease on:**

**Renal Drug Elimination**

**Hepatic Drug Metabolism**

**Drug Transporters**

**Drug Distribution**

**Drug Absorption**

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# GOALS Of Effects of Renal Disease on PK Lecture

- ***DOSE ADJUSTMENT* in Patients with Renal Impairment**

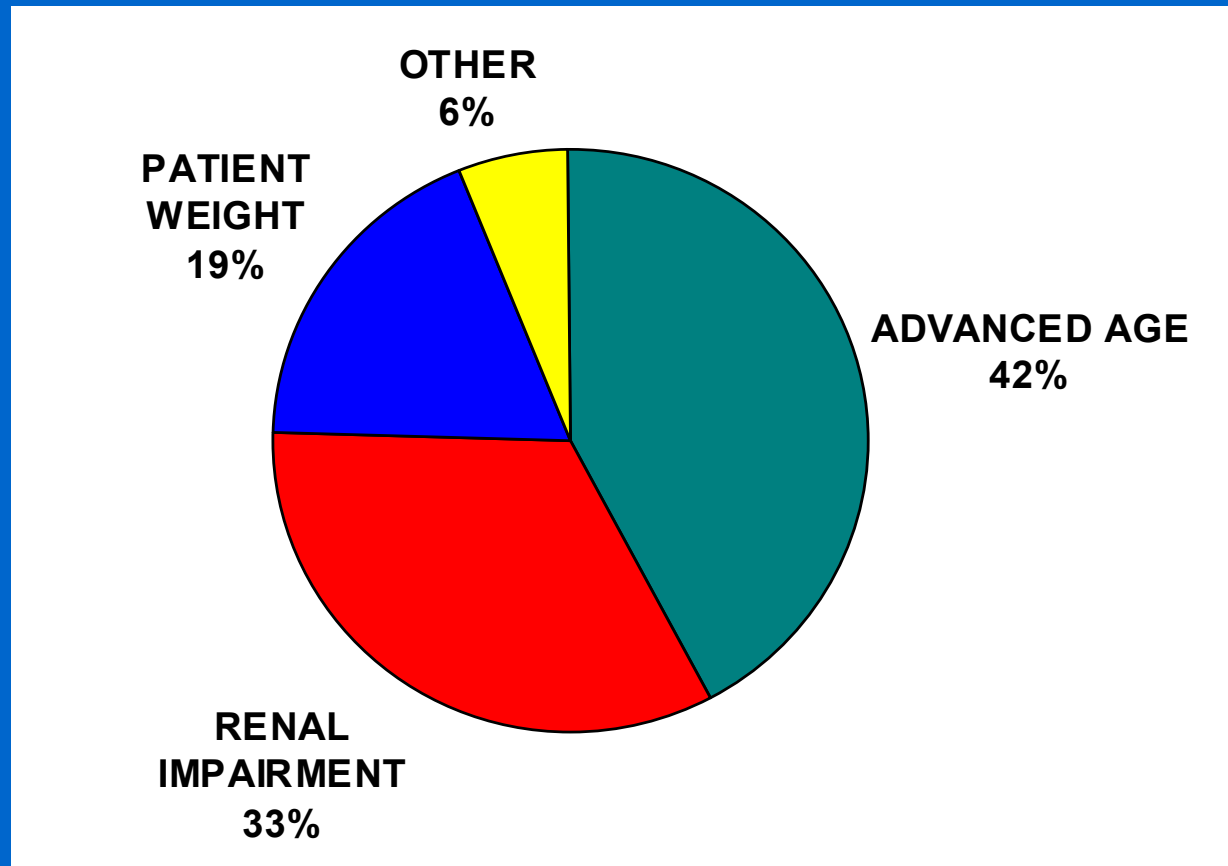
## **Statement of the Problem**

**How is renal function assessed?**

**How is drug dose adjusted based on this assessment?**

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# *PATHOPHYSIOLOGIC FACTORS NOT ACCOUNTED FOR IN DRUG DOSING\**



\* Lesar TS, Briceland L, Stein DS. JAMA 1997;277:312-7.

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# ***INFORMATION CONTENT OF CURRENT DRUG LABELS\****

<b>CORE INFORMATION CATEGORY</b>	<b>Inclusion of Desirable Data Elements MEAN (95% CI)</b>	
<i><b>MECHANISM OF ACTION</b></i>	<b>88%</b>	<b>(84% - 93%)</b>
<i><b>PHARMACODYNAMICS</b></i>	<b>43%</b>	<b>(37% - 49%)</b>
<i><b>DRUG METABOLISM</b></i>	<b>23%</b>	<b>(16% - 29%)</b>
<i><b>PHARMACOKINETICS</b></i>	<b>42%</b>	<b>(35% - 49%)</b>
<i><b>DOSE ADJUSTMENT</b></i>	<b>37%</b>	<b>(32% - 42%)</b>

\* Spyker DA, et al. Clin Pharmacol Ther 2000;67:196-200.

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# ***FDA GUIDANCE FOR INDUSTRY***

***PHARMACOKINETICS IN PATIENTS WITH  
IMPAIRED RENAL FUNCTION*** – Study  
Design, Data Analysis, and Impact on Dosing  
and Labeling (1998)

**AVAILABLE AT:**

**<http://www.fda.gov/cder/guidance/index.htm>**

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# GOALS of Renal Disease Effects Lecture

- ***DOSE ADJUSTMENT* in Patients with Renal Impairment**
  - Statement of the Problem
  - **How is renal function assessed?**
  - How is drug dose adjusted based on this assessment?



# *ELIMINATION* by Different Routes

MEASUREMENTS	RENAL	HEPATIC	DIALYSIS
Blood Flow	+	+	+
Afferent Concentration	+	+	+
Efferent Concentration	0	0	+
Eliminated Drug	+	0	+

*\*not actually measured in routine PK studies*

# *RENAL CLEARANCE* EQUATION

$$CL = \frac{U \times V}{P}$$

**U = URINE CONCENTRATION**

**V = URINE VOLUME / TIME**

**P = PLASMA CONCENTRATION**

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## ***CLEARANCE TECHNIQUES FOR ASSESSING RENAL FUNCTION***

### **GLOMERULAR FILTRATION:**

Normal: 120 – 130 mL/min/1.73 m<sup>2</sup>

#### ***CLEARANCE MARKERS:***

Inulin

Creatinine

<sup>125</sup>I-Iothalamate

### **RENAL BLOOD FLOW:**

Normal: ♂ 1,209 ± 256 mL/min/1.73 m<sup>2</sup>

♀ 982 ± 184 mL/min/1.73 m<sup>2</sup>

#### ***CLEARANCE MARKER:***

Para-Aminohippuric Acid

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## GOALS of Renal Disease Effects Lecture

### - How is renal function assessed?

Commonly estimated from the *Cockcroft and Gault equation* for creatinine clearance if renal function is stable, but the *Modification of Diet in Renal Disease (MDRD) Study equation* for estimating GFR is now the preferred approach.

# Estimation of GFR

- The **MDRD equation** to estimate GFR from serum creatinine is **the most accurate** compared to the (125)I-iothalamate standard.
- However, it tends to underestimate high GFRs and also overestimates low GFRs.

Levey AS et al. *Ann Intern Med.* 2006;145-247-254

[http://www.nkdep.nih.gov/professionals/gfr\\_calculators/idms\\_con.htm](http://www.nkdep.nih.gov/professionals/gfr_calculators/idms_con.htm)

# Renal Clearance of Drugs

- Generally, there is a **linear correlation** between the clearance of creatinine and the clearance of drugs excreted via the kidneys.
- We take advantage of this correlation when making **dose adjustments** in patients with impaired renal function.

# *STEADY STATE* CONCENTRATION

## Continuous Infusion:

$$C_{SS} = \frac{I}{CL_E}$$

## Intermittent Dosing:

$$\bar{C}_{SS} = \frac{DOSE/\tau}{CL_E}$$

# *ADDITIVITY* OF CLEARANCES

$$CL_E = CL_R + CL_{NR}$$

$CL_R$  = RENAL CLEARANCE

$CL_{NR}$  = NON-RENAL CLEARANCE



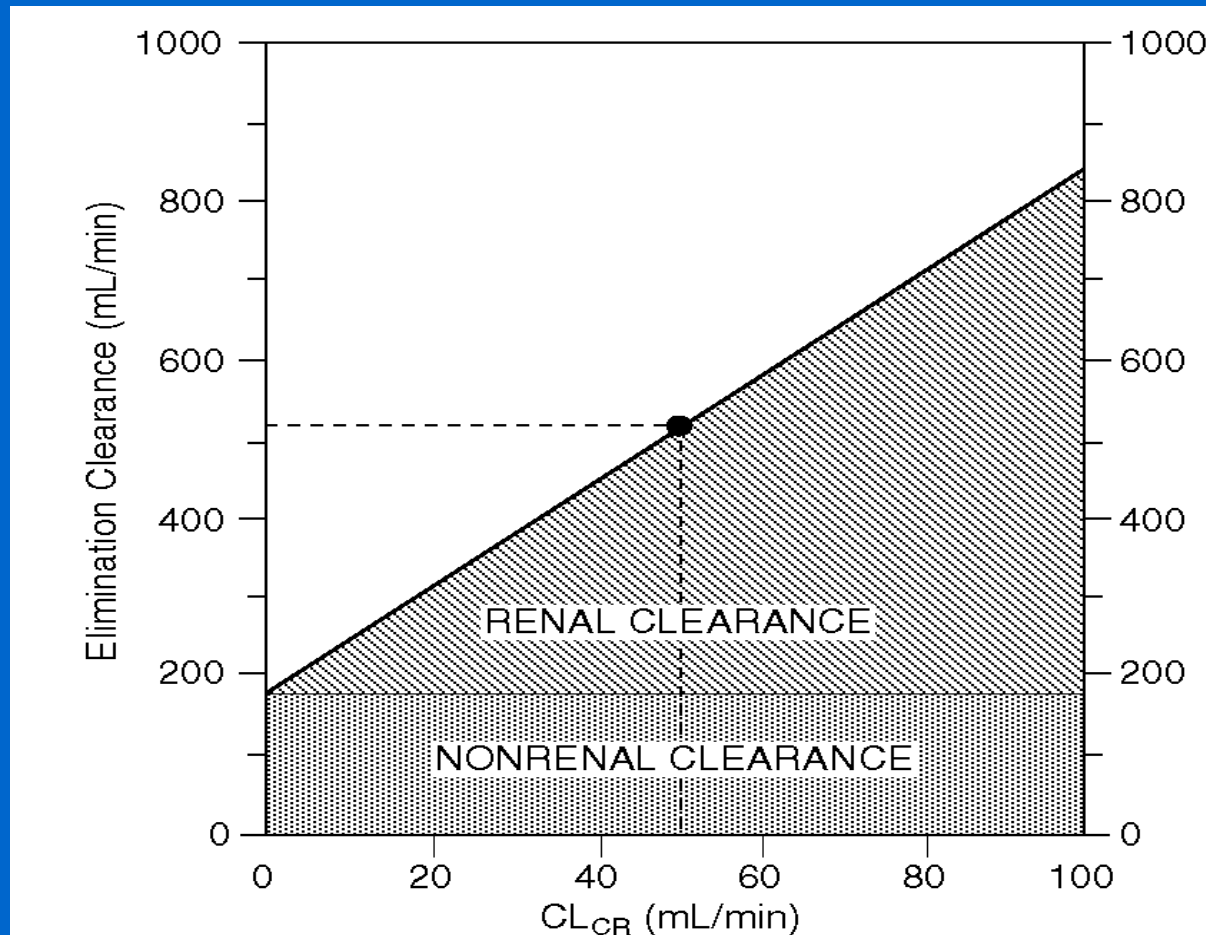
# DETTLI Approach\*

$$CL_R = \alpha CL_{Cr}$$

$$CL_E = CL_R + CL_{NR}$$

\* Dettli L. Med Clin North Am 1974;58:977-85

# *NOMOGRAM* FOR **CIMETIDINE** DOSING\*



\*From: Atkinson AJ Jr, Craig RM. Therapy of peptic ulcer disease.

# Key *ASSUMPTIONS* of Dettli Method

- $CL_{NR}$  remains *CONSTANT* when renal function is impaired.
- $CL_R$  declines in *LINEAR FASHION* with  $CL_{CR}$ 
  - *Intact Nephron Hypothesis*
  - Some drugs  $\downarrow$  *SECRETION*  $>$  *GFR* with aging\*

\* Reidenberg MM, et al. Clin Pharmacol Ther 1980;28:732-5.

# CIMETIDINE Case History

A 67-year-old veteran had been **functionally anephric**, requiring outpatient **hemodialysis** for several years. He was hospitalized for revision of his arteriovenous shunt and postoperatively complained of symptoms of **gastroesophageal reflux**. This complaint prompted institution of **cimetidine** therapy in a dose of 300 mg every 6 hours.

## CIMETIDINE Case History (cont.)

### Rationale for Prescribed Cimetidine Dose:

*At that time, 600 mg every 6 hours was the usual cimetidine dose for patients with normal renal function and the **Physician's Desk Reference** recommended halving the cimetidine dose for patients “with creatinine clearance less than 30 cc/min”.*

## CIMETIDINE Case History (cont.)

Three days later the patient was noted to be **confused**. The nephrology team reevaluated the patient and agreed to *discontinue cimetidine* as suggested by the attending internist/clinical pharmacologist. Two days later the patient was **alert** and was discharged from the hospital to resume outpatient hemodialysis therapy.

## ***LABELING*** FOR CIMETIDINE\*

- ***DOSAGE ADJUSTMENT***

1/2 normal dose if  $CL_{Cr} < 30$  mL/min

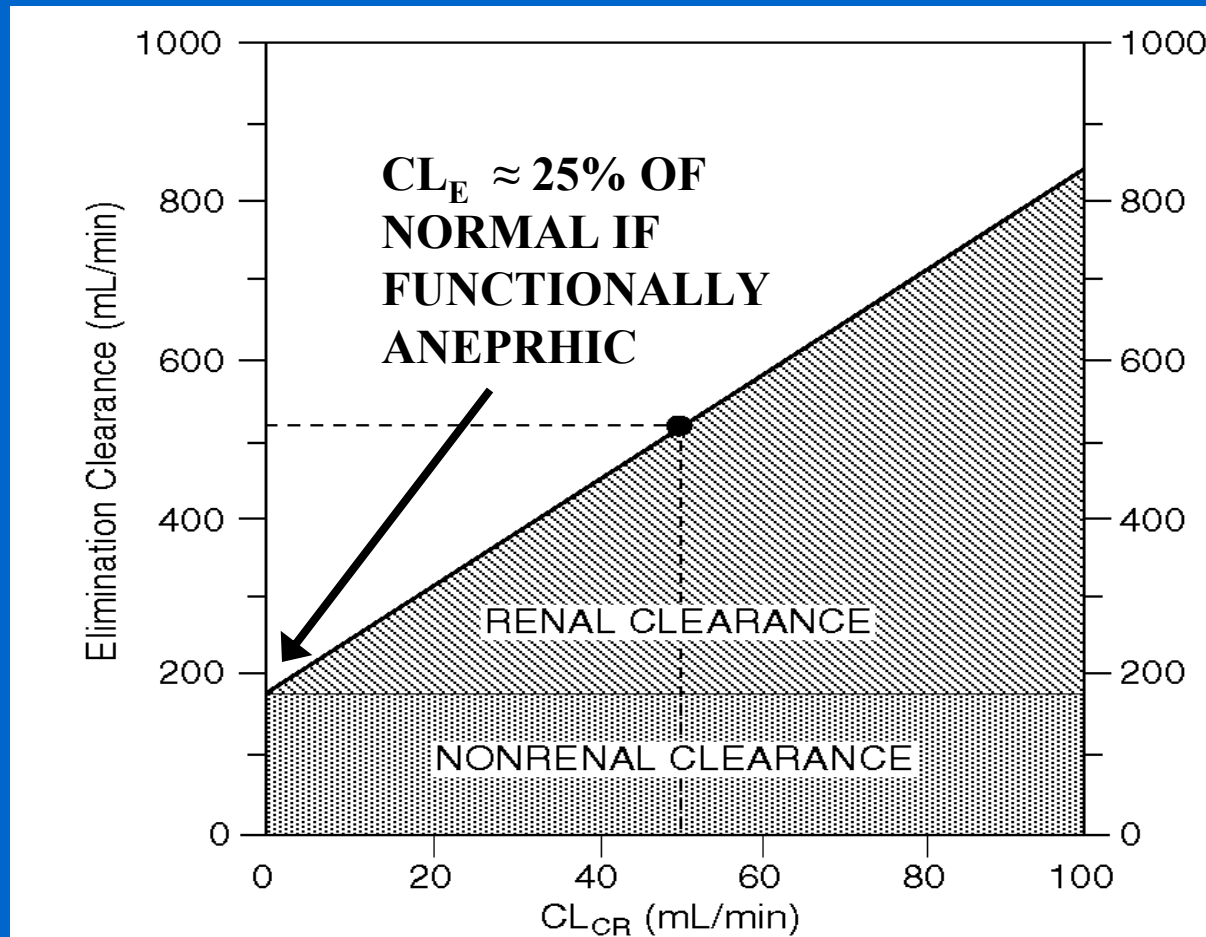
- ***PHARMACOKINETICS***

Following I.V. or I.M. administration in *normal subjects*,

**~ 75% of drug is recovered from the urine as parent compound.**

\* Physician's Desk Reference. 58<sup>th</sup> edition, 2004.

# *NOMOGRAM* FOR **CIMETIDINE** DOSING\*



\*From: Atkinson AJ Jr, Craig RM. Therapy of peptic ulcer disease.



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## ***DOSE ADJUSTMENT OPTIONS FOR PATIENTS WITH RENAL IMPAIRMENT***

$$\bar{C}_{SS} = \frac{\text{DOSE} / \tau}{\text{CL}_E}$$

- MAINTAIN USUAL DOSING INTERVAL BUT ***REDUCE DOSE*** IN PROPORTION TO  $\downarrow \text{CL}_E$
  - MAINTAIN USUAL DOSE BUT ***INCREASE DOSING INTERVAL*** IN PROPORTION TO  $\downarrow \text{CL}_E$
  - ***ADJUST BOTH*** DOSE AND DOSING INTERVAL
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# GOALS of Renal Disease Effects Lecture

- **EFFECT OF RENAL DISEASE ON RENAL DRUG ELIMINATION**
  - *MECHANISMS* OF RENAL DRUG ELIMINATION
  - CONCEPT OF *RESTRICTIVE VS. NONRESTRICTIVE* ELIMINATION

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# ***MECHANISMS* of Renal Drug Elimination**

**Glomerular Filtration**

**Renal Tubular Secretion**

**Reabsorption by Non-Ionic Diffusion**

**Active Reabsorption**

# MECHANISMS OF RENAL ELIMINATION

## GLOMERULAR FILTRATION

- Affects all drugs and metabolites of **appropriate molecular size**.
- *Influenced* by **protein binding**

$$\text{Drug Filtration Rate} = \text{GFR} \times f_u \times [\text{Drug}]$$

( $f_u$  = free fraction)

## RENAL TUBULAR SECRETION

- *Not influenced* by protein binding
- May be affected by *other drugs*, etc.

### EXAMPLES:

Active Drugs:      ACIDS – Penicillin  
                             BASES – Procainamide

Metabolites:        Glucuronides, Hippurates, etc.

# *RESTRICTIVE vs. NONRESTRICTIVE ELIMINATION*

## **RESTRICTIVE:**

Clearance *DEPENDS* on Protein Binding.

KIDNEY: Drug Filtration Rate =  $f_u \cdot \text{GFR}$

LIVER:  $\text{CL} = f_u \cdot \text{Cl}_{\text{int}}$

## **NONRESTRICTIVE:**

Clearance *INDEPENDENT* of Protein Binding

KIDNEY:  $\text{CL} = Q$  (renal blood flow)

***EXAMPLE: PARA-AMINOHIPPURATE CLEARANCE  
MEASURES RENAL BLOOD FLOW.***

# INTRINSIC CLEARANCE

*INTRINSIC CLEARANCE* IS THE  
ELIMINATION CLEARANCE THAT  
*WOULD BE OBSERVED* IN THE  
*ABSENCE OF ANY PROTEIN BINDING*  
*RESTRICTIONS.*

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# *RESTRICTIVE vs. NONRESTRICTIVE ELIMINATION*

## RESTRICTIVE:

Clearance *DEPENDS* on Protein Binding

KIDNEY: Drug Filtration Rate =  $f_u \cdot \text{GFR}$

LIVER:  $\text{CL} = f_u \cdot \text{Cl}_{\text{int}}$

## NONRESTRICTIVE:

Clearance *INDEPENDENT* of Protein Binding

KIDNEY:  $\text{CL} = Q$  (renal blood flow)

LIVER:  $\text{CL} = Q$  (hepatic blood flow)

# Renal *REABSORPTION* Mechanisms

## REABSORPTION BY NON-IONIC DIFFUSION

- Affects **weak acids** and **weak bases**.
- Only important if excretion of *free drug* is major elimination pathway.

### *EXAMPLES:*

Weak Acids:	PHENOBARBITAL
Weak Bases:	QUINIDINE

## ACTIVE REABSORPTION

- Affects **ions**, not proved for other drugs.

### *EXAMPLES:*

Halides:	FLUORIDE, BROMIDE
Alkaline Metals:	LITHIUM



## *RENAL EXCRETION* OF DRUGS

INTACT NEPHRON HYPOTHESIS: Provides a basis for dose adjustment when renal excretion of drug is impaired.

- Regardless of mechanism, *renal drug elimination declines in parallel with decreases in GFR.*
- Therefore,  $CL_{Cr}$  can be used to assess impact of renal impairment on renal excretion of drugs.

WHAT ABOUT OTHER EXCRETION ROUTES?

# GOALS of Renal Disease Effects Lecture

- EFFECT OF RENAL DISEASE ON *DRUG METABOLISM and TRANSPORT*

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# CRF – Effects on Drug Metabolism and Transport

Recent **Reviews** on this topic:

TD Nolin, J Naud, FA Leblond, V Pichette

Emerging Evidence of the Impact of  
Kidney Disease on Drug Metabolism  
and Transport

*Clin. Pharmacol. Ther.* 2008;83:898-903

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# CRF – Effects on Drug Metabolism and Transport

Recent **Reviews** on this topic:

AW Dreisbach, JJJ Lertora

The effect of chronic renal failure on  
drug metabolism and transport

*Expert Opin. Drug Metab. Toxicol.*

*2008;4:1065-1074*

⋮

# Effect of CRF on Non-Renal Drug Clearance in Humans

	<b>CL<sub>NR</sub> (%)</b>	<b>Enzyme</b>
<b>Captopril</b>	<b>- 50</b>	<b>TPMT</b>
<b>Morphine</b>	<b>- 40</b>	<b>UGT2B7</b>
<b>Procainamide</b>	<b>- 60</b>	<b>NAT-2</b>
<b>Verapamil</b>	<b>- 54</b>	<b>CYP3A4</b>
<b>Metoclopramide</b>	<b>- 66</b>	<b>CYP2D6</b>
<b>Warfarin</b>	<b>- 50</b>	<b>CYP2C9</b>

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# Effect of CRF on Drug Transport

**Impaired transport function in renal failure (intestine, liver, kidney)**

- **P-Glycoprotein**
- **Organic Anion Transporting Polypeptide (OATP)**

*Fexofenadine is a substrate for both*

# Effect of CRF on Bioavailability

Studies in human subjects:

Propranolol	+300 %	CYP2D6
Erythromycin	+100 %	CYP3A4
Propoxyphene	+100 %	CYP3A4
Dyhydrocodeine	+70 %	CYP2D6

# Effects of Uremic Toxins

Indoxyl sulfate

CMPF-propanoic acid

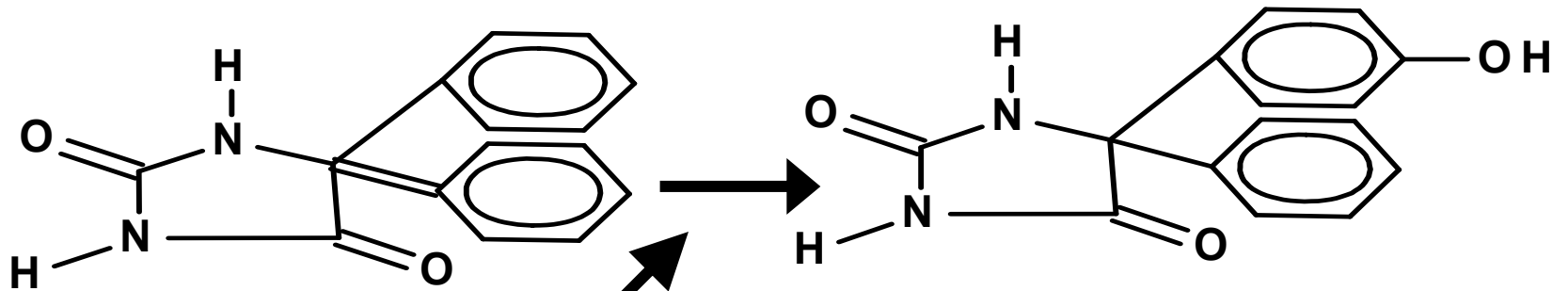
Parathyroid hormone (PTH)

Cytokines (chronic inflammation)

Inhibition of drug metabolism and  
transport **reversed by hemodialysis**



# PHASE I AND PHASE II METABOLIC REACTIONS

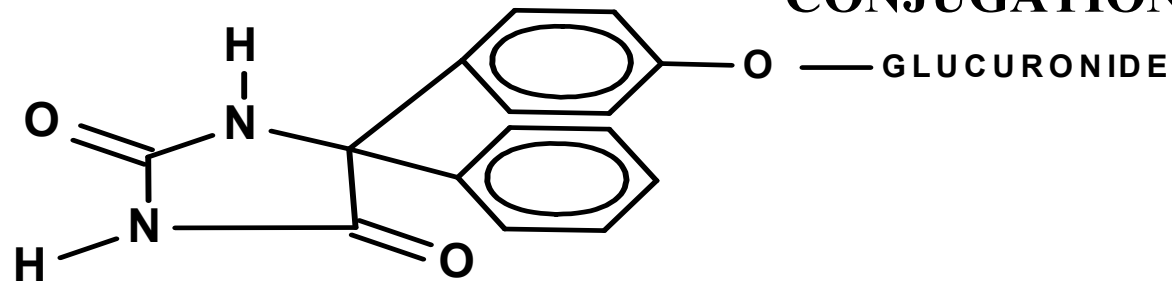


PHENYTOIN

*p* - HPPH

**PHASE I**  
**HYDROXYLATION**

**PHASE II**  
**GLUCURONIDE**  
**CONJUGATION**

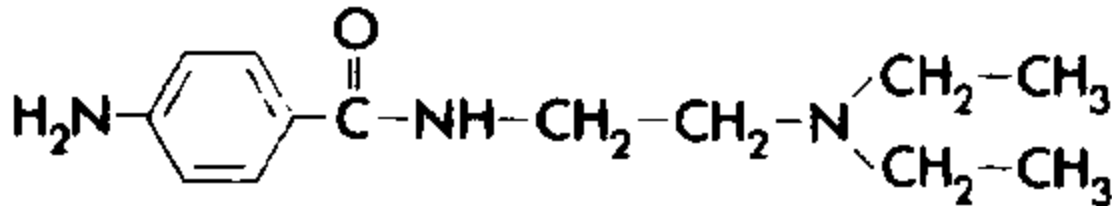


*p* - HPPH GLUCURONIDE

# GOALS of Renal Disease Effects Lecture

- EFFECT OF RENAL DISEASE ON *DRUG METABOLISM*
- *EXAMPLES:*
  - PROCAINAMIDE** - Acetylation
  - PHENYTOIN** - Hydroxylation

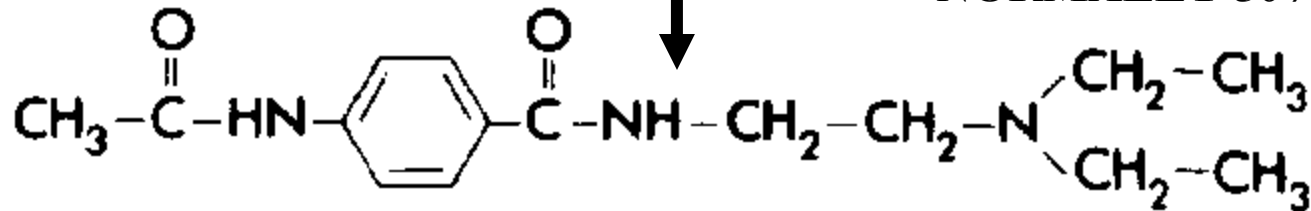
# PROCAINAMIDE ACETYLATION



**PROCAINAMIDE**

NAT2: FAST VS. SLOW

RENAL ELIMINATION NORMALLY 50%



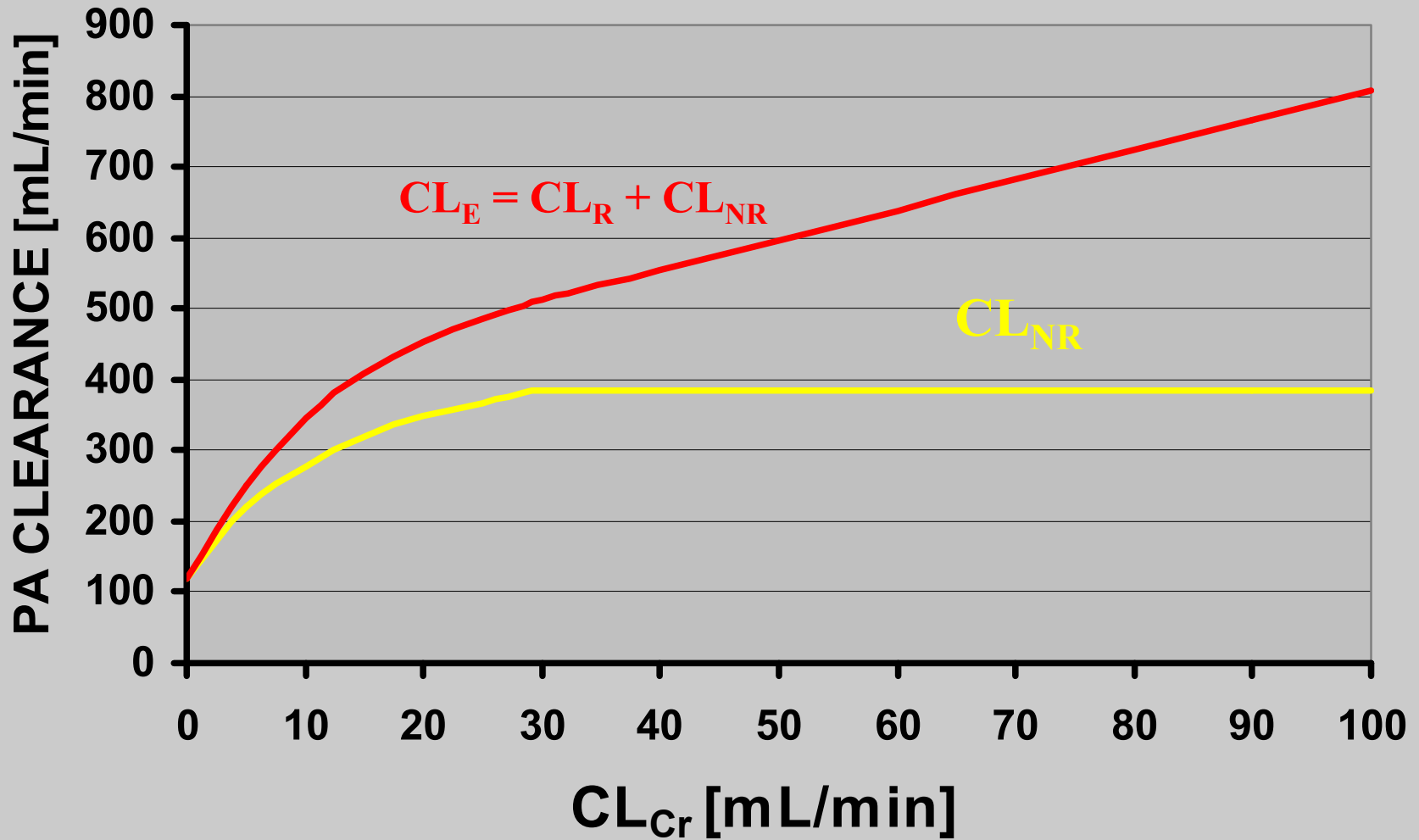
**N-ACETYLPROCAINAMIDE (NAPA)**

# Procainamide Kinetics in *DIALYSIS PATIENTS\**

	<i>NORMALS</i>		<i>FUNCTIONALLY ANEPHRIC PATIENTS</i>	
	Fast	Slow	Fast	Slow
$T_{1/2}$ (hr)	2.6	3.5	12.2	17.0
$CL_E$ (L/kg)	809	600	118	94
$CL_R$ (L/kg)	426	357	0	0
$CL_{NR}$ (L/kg)	383	243	118	94
$V_{d(ss)}$ (L/kg)	1.95	1.93	1.41	1.93

\* From: Gibson TP. *Kidney Int* 1977;12:422-9.

# Procainamide Dosing Nomogram (FAST ACETYLATORS)



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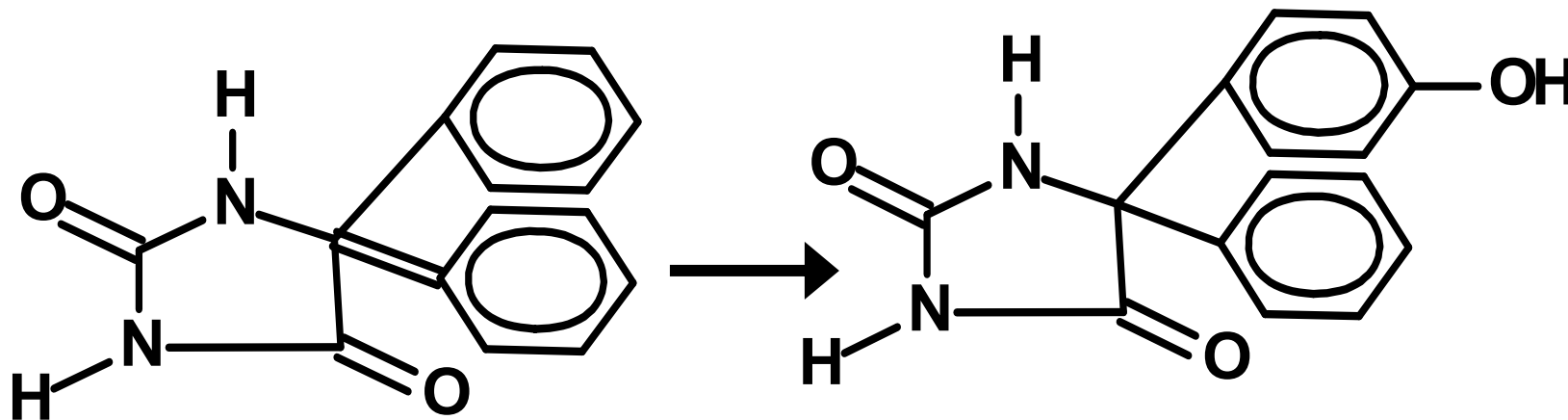
## **NAPA** ELIMINATION HALF LIFE IN *FUNCTIONALLY ANEPHRIC PATIENTS*

- **HEALTHY SUBJECTS:** 6.2 hr
- ***PREDICTED*** for DIALYSIS PATIENTS: 42.8 hr \*
- ***MEASURED*** in DIALYSIS PATIENTS: 41.9 hr \*

\* See Study Problem at end of Chapter 5.

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# PHENYTOIN *HYDROXYLATION* BY P450

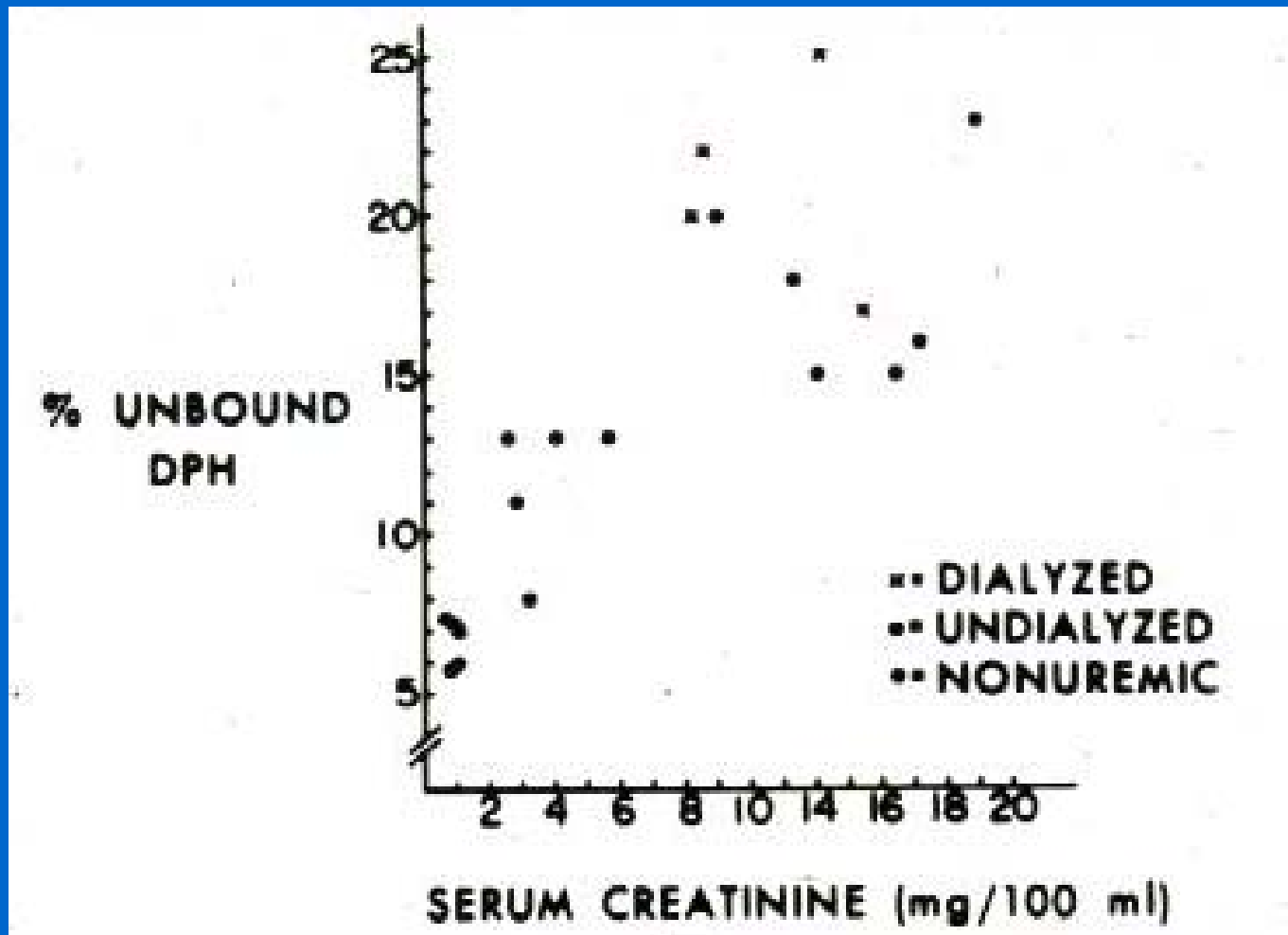


PHENYTOIN

*p* - HPPH

CYP2C9: Major, CYP2C19: Minor

# Effect of Renal Disease on *PHENYTOIN* *PROTEIN BINDING*





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# PHENYTOIN

## *KINETICS IN DIALYSIS PATIENTS\**

	NORMALS (N = 4)	UREMIC PATIENTS (N = 4)
<b>% UNBOUND (<math>f_u</math>)</b>	<b>12%</b>	<b>26%</b>
$CL_H$	2.46 L/hr	7.63 L/hr
$CL_{int}$	20.3 L/hr	29.9 L/hr <b>NS</b>

$$CL_H = f_u \cdot Cl_{int}, \quad \text{So: } Cl_{int} = CL_H / f_u$$

\* From: Odar-Cederlöf I, Borgå O: Eur J Clin Pharmacol 1974;7:31-7.

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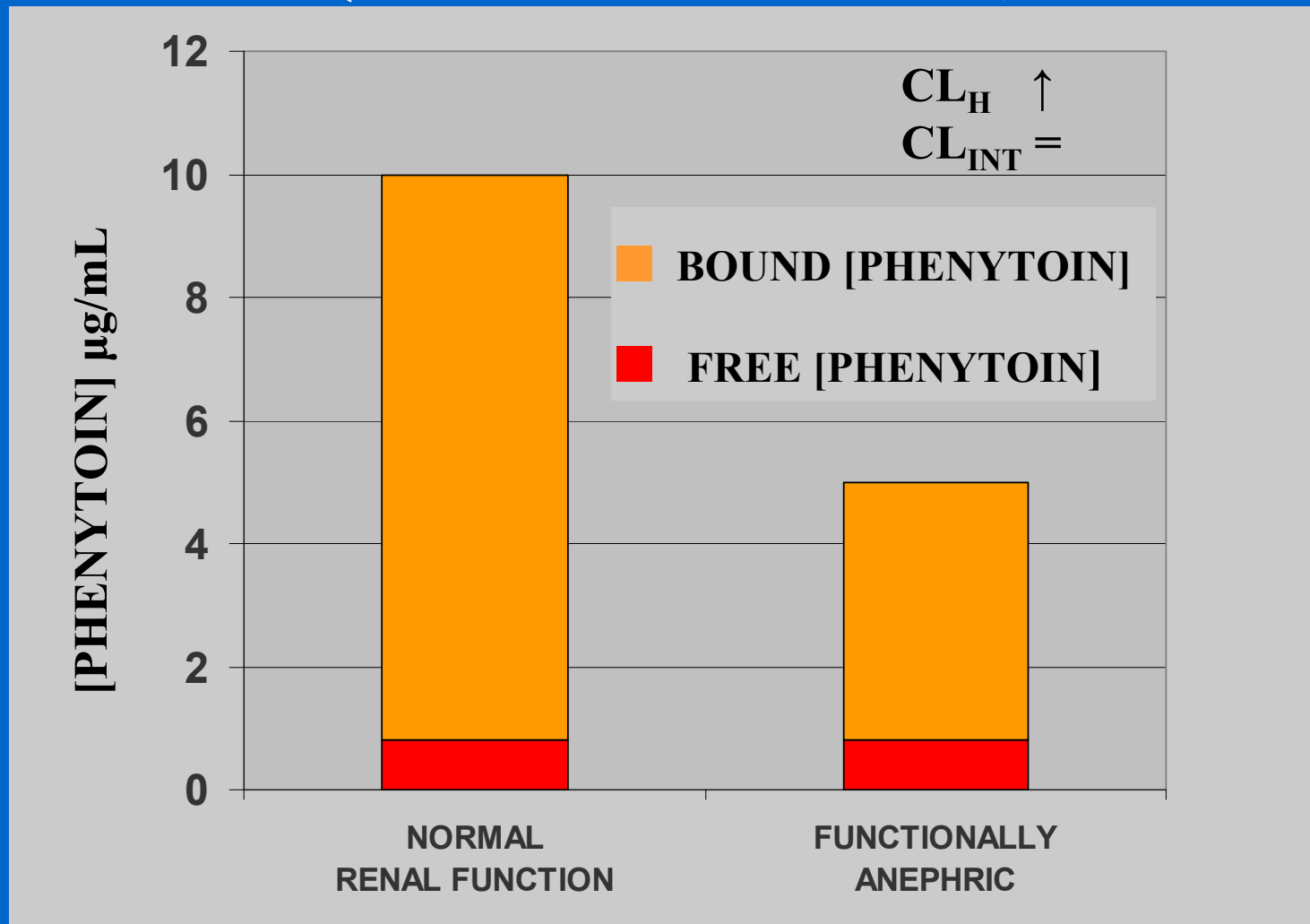
## Effect of *PROTEIN BINDING* Changes on **Phenytoin** Plasma Concentration

$$\bar{C}_{ss} = \frac{\text{DOSE} / \tau}{\text{CL}_E}$$

**PHENYTOIN > 98% ELIMINATED BY HEPATIC METABOLISM, SO  $\text{CL}_E = \text{CL}_H$**

$$\bar{C}_{ss, u} / f_u = \frac{\text{DOSE} / \tau}{f_u \text{CL}_{INT}}$$

# ***FREE AND TOTAL PHENYTOIN LEVELS*** **(DOSE = 300 MG/DAY)**



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# ***THERAPEUTIC RANGE*** of **Phenytoin** Levels in *Dialysis Patients*

*RISK is that **TOTAL** levels below the usual range of 10 – 20 µg/mL will prompt inappropriate dose adjustment in dialysis patients.*      ↑

## **THERAPEUTIC RANGE FOR DIALYSIS PTS:**

**Based on “Total Levels”:**      5 - 10 µg/mL

**Based on “Free Levels”:**      0.8 - 1.6 µg/mL

# GOALS of Renal Disease Effects Lecture

- **EFFECT OF RENAL DISEASE ON DRUG DISTRIBUTION**

- **PLASMA PROTEIN BINDING**

*EXAMPLE: PHENYTOIN*

- **TISSUE BINDING**

*EXAMPLE: DIGOXIN*

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## Effect of Renal Disease on *BINDING TO PLASMA PROTEINS\**

*BASIC OR NEUTRAL  
DRUGS:*

NORMAL OR  
SLIGHTLY REDUCED

*ACIDIC DRUGS:*

REDUCED FOR MOST

\* From: Reidenberg MM, Drayer DE: Clin Pharmacokinet  
1984;9(Suppl. 1):18-26.

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## Effect of Binding Changes on *APPARENT DISTRIBUTION VOLUME\**

$$V_d = ECF + \phi f_u (TBW - ECF)$$

$\Phi$  = TISSUE/PLASMA PARTITION RATIO

$f_u$  = FRACTION NOT BOUND TO PLASMA  
PROTEINS

FOR PHENYTOIN:  $\Phi = 10.4$

\* Atkinson AJ Jr, et al. Trends Pharmacol Sci 1991;12:96-101.

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# PHENYTOIN *DISTRIBUTION* IN *DIALYSIS* PATIENTS\*

	NORMALS	UREMIC PATIENTS
% UNBOUND ( $f_u$ )	12% <sup>†</sup>	26%
$V_{d(AREA)}$	0.64 L/kg	1.40 L/kg

<sup>†</sup> USUAL VALUE IN NORMAL SUBJECTS ~ 9%

\* From: Odar-Cederlöf I, Borgå O: Eur J Clin Pharmacol 1974;7:31-7.

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# GOALS OF RENAL DISEASE EFFECTS LECTURE

- **EFFECT OF RENAL DISEASE ON DRUG  
DISTRIBUTION**

- **PLASMA PROTEIN BINDING**

*EXAMPLE: PHENYTOIN*

- **TISSUE BINDING**

*EXAMPLE: DIGOXIN*

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**IMPAIRED RENAL FUNCTION *REDUCES*  
DIGOXIN *DISTRIBUTION VOLUME\****

$$V_d = 3.84 \cdot \text{wt (kg)} + 3.12 \text{ CL}_{\text{cr}} \text{ (mL/min)}$$

\* Sheiner LB, et al. J Pharmacokinet Biopharm 1977;5:445-79.

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# ***CRITERIA FOR NORMAL ABSORPTION OF 25 GRAM D-XYLOSE DOSE***

<b>5-hr URINE RECOVERY</b>	<b>&gt; 4 g</b>
<b>[SERUM] 1 hr AFTER DOSE</b>	<b>≥ 0.2 mg/mL</b>
<b>% DOSE ABSORBED</b>	<b>&gt; 42%</b>
<b><math>k_a</math></b>	<b>&gt; 0.37 hr<sup>-1</sup></b>

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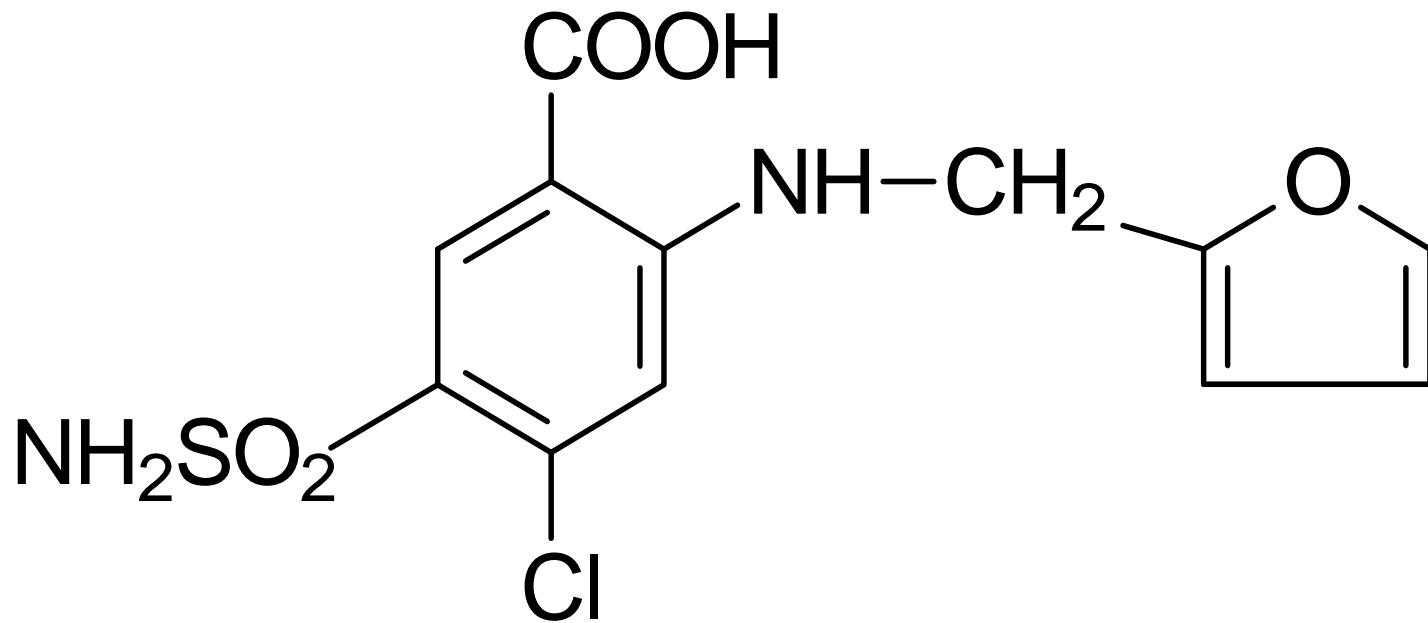
## EFFECT OF RENAL DISEASE ON **D-XYLOSE** *ABSORPTION*\*

PATIENT GROUP	$k_a$ (hr <sup>-1</sup> )	$k_o$ (hr <sup>-1</sup> )	% DOSE ABSORBED
NORMALS	<b>1.03 ± 0.33</b>	0.49 ± 0.35	<b>69.4 ± 13.6</b>
MODERATE	0.64 ± 0.28	0.19 ± 0.15	77.4 ± 14.8
DIALYSIS	<b>0.56 ± 0.42</b>	0.67 ± 0.61	<b>48.6 ± 13.3</b>

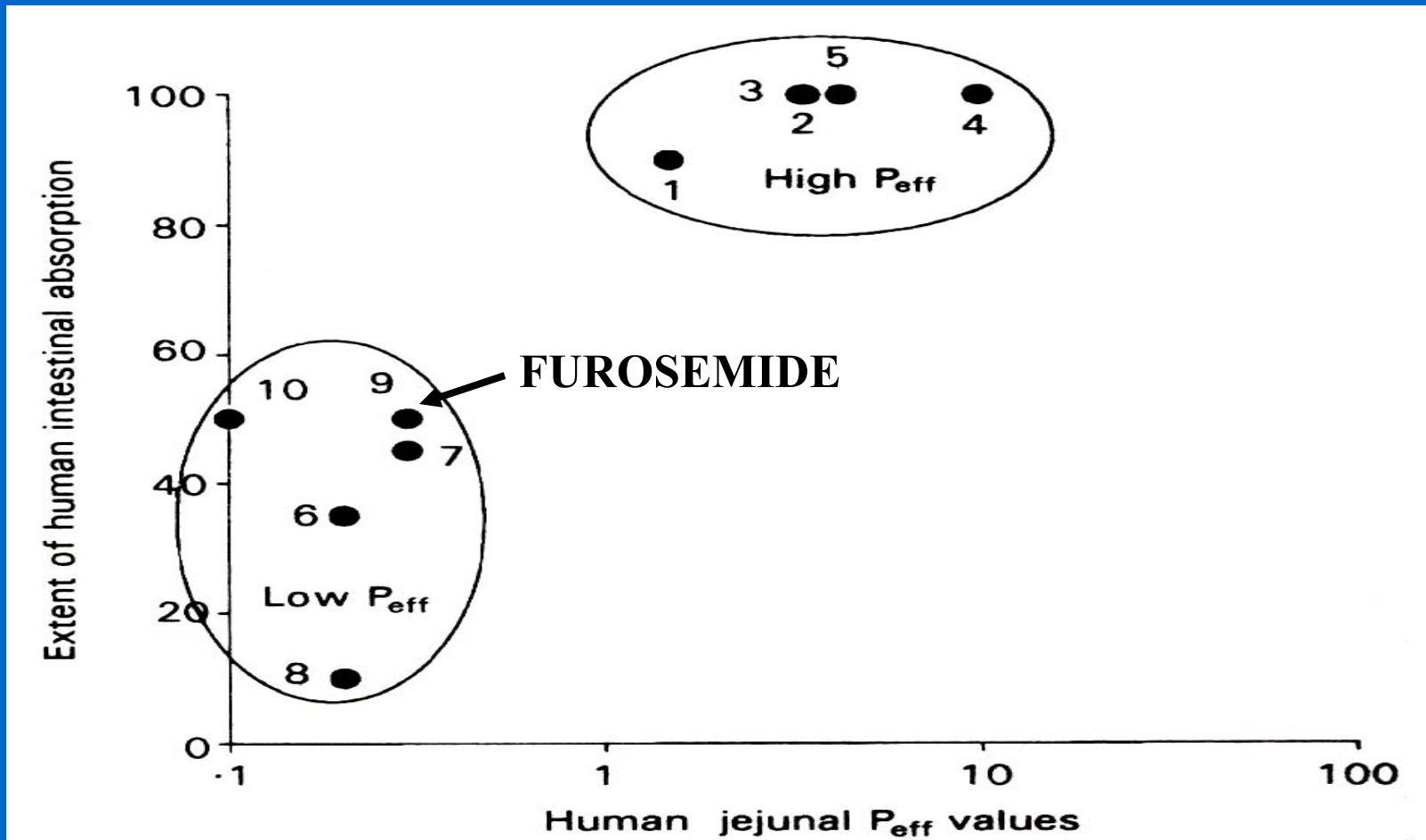
\* From: Worwag EM et al. Clin Pharmacol Ther 1987;41:351-7.

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# FUROSEMIDE



# BIOPHARMACEUTIC CLASSIFICATION OF FUROSEMIDE\*



\* From: Lenneräs. J Pharm Pharmacol 1997;49:627-38.

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# BIOPHARMACEUTIC DRUG CLASSIFICATION OF **FUROSEMIDE** \*

CLASS IV:

LOW SOLUBILITY-LOW PERMEABILITY

- *in vitro* – *in vivo* correlation poor
- good bioavailability not expected

\* From: Lenneräs, et al. Pharm Res 1995;12:S396

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# Biopharmaceuticals Classification System (BCS)

- **Class I (high S, high P)**  
*Enzyme effects predominate*
- **Class II (low S, high P)**  
*Both enzymes and transporters*
- **Class III (high S, low P)**  
*Transporter effects predominate*

Sun H, et al (2006)

Amidon GI, et al (1995)



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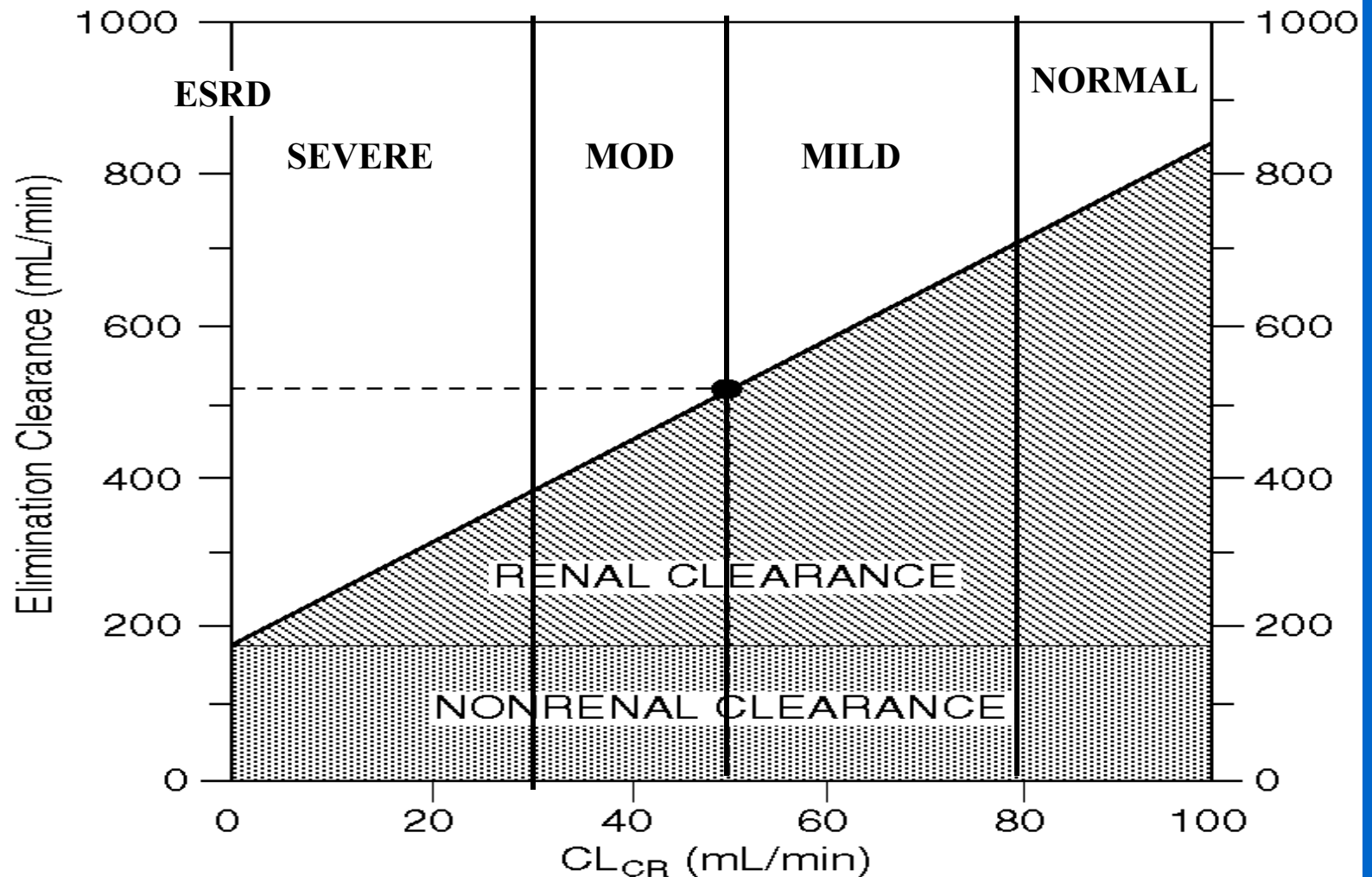
# ***FDA GUIDANCE FOR INDUSTRY***

## ***PHARMACOKINETICS IN PATIENTS WITH IMPAIRED RENAL FUNCTION*** – Study Design, Data Analysis, and Impact on Dosing and Labeling

**AVAILABLE AT:**

**<http://www.fda.gov/cder/guidance/index.htm>**

# BASIC “FULL” STUDY DESIGN



# Effects of Hemodialysis

**Advanced CRF:**

**Stage IV (GFR 15-29 ml/min)**

**Stage V (GFR 0-15 ml/min)**

**Hemodialysis** may reverse the inhibition of drug metabolizing enzymes and transporters

## FDA *GUIDANCE FOR INDUSTRY*

- A **revision** of this guidance document is currently under way (2008).
- A **concept paper/draft guidance** has been posted by the FDA regarding revised recommendations for PK studies in patients with **impaired renal function**.

<http://www.fda.gov/ohrms/dockets/ac/08/briefing/2008-4351b1-01-FDA.pdf>

(document pages 57-73)