

# PHARMACOKINETICS IN PATIENTS REQUIRING RENAL REPLACEMENT Rx

## PART 1: PK IN PATIENTS REQUIRING HEMODIALYSIS

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## JOHN JACOB ABEL 1857 - 1938



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## FIRST DESCRIPTION OF HEMODIALYSIS IN ANIMALS\*

### ON THE REMOVAL OF DIFFUSIBLE SUBSTANCES FROM THE CIRCULATING BLOOD OF LIVING ANIMALS BY DIALYSIS

JOHN J. ABEL, LEONARD G. BOWNTREE AND B. B. TURNER  
From the Pharmacological Laboratory of the Johns Hopkins University

Received for publication, December 18, 1913

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\* From: Abel JJ, et al. J Pharmacol Exp Ther 1914;5:275-317.

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## WILLEM J. KOLFF, M.D. (1911 - )



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## ELIMINATION BY DIFFERENT ROUTES

MEASUREMENTS	RENAL	HEPATIC	DIALYSIS
BLOOD FLOW	+*	+*	+
AFFERENT CONC.	+	+	+
EFFERENT CONC.	0	0	+
ELIMINATED DRUG	+	0	+

\*not actually measured in routine PK studies

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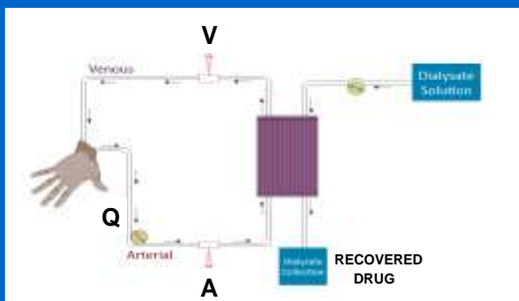
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## DATA SOURCES FOR FICK EQUATION



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## IMPACT OF $CL_D$

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

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## CRITERION FOR DIALYSIS EFFICACY\*

$$CL_{EC} > 30\% [CL_R + CL_{NR}]$$

**BUT CLEARANCE ESTIMATES  
MUST BE COMPARABLE**

\* Levy G. Am J Med 1977;62:461-5.

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## GOALS OF DIALYSIS DISCUSSION

DISCUSSION OF DIALYSIS CLEARANCE  
MECHANISTIC - RENKIN APPROACH  
EMPIRICAL  
RECOVERY CLEARANCE  
FICK EQUATION  
CLINICAL STUDIES OF DIALYSIS PK  
MODEL PROSPECTIVE STUDY  
TREATMENT OF DRUG TOXICITY  
PHYSIOLOGIC CHANGES DURING DIALYSIS  
USE OF KINETIC METHODS FOR ANALYSIS  
PATHOPHYSIOLOGIC CONSEQUENCES

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**EUGENE RENKIN  
PROFESSOR EMERITUS AT UC DAVIS**



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**RENKIN DIALYSIS EQUATION\***

$$CL_D = Q(1 - e^{-P \cdot S / Q})$$

**Q** = DIALYZER BLOOD FLOW

**P·S** = PERMEABILITY-SURFACE AREA  
PRODUCT OF DIALYZING MEMBRANE

**NEGLECTS: BOUNDARY EFFECTS, ULTRAFILTRATION**

\* From Renkin EM. Tr Am Soc Artific Organs 1956;2:102-5

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**DETERMINANTS OF  
PERMEABILITY TERM (P or P · S)**

- DIALYZER MEMBRANE CHARACTERISTICS
  - MEMBRANE SURFACE AREA
  - MEMBRANE THICKNESS
  - MEMBRANE POROSITY
- DRUG BINDING TO PLASMA PROTEINS
- SOLUTE SIZE AND DIFFUSIVITY

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## DIALYZER PERMEABILITY VS. FREE WATER DIFFUSION COEFFICIENTS

### PROCAINAMIDE/NAPA:

RATIO OF DIALYZER PERMEABILITY COEFFICIENTS\*  $1.28 \pm 0.23$

RATIO OF FREE WATER DIFFUSION COEFFICIENTS 1.23

\* From Gibson TP et al. Clin Pharmacol Ther 1976;20:720-6.

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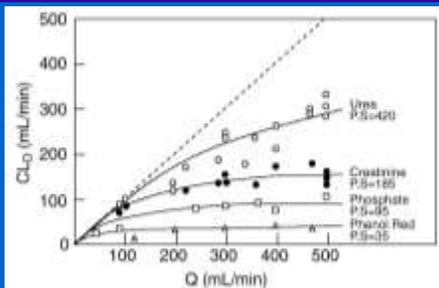
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## DIALYSIS CLEARANCE VS. DIALYZER BLOOD FLOW\*



\* From Renkin EM. Tr Am Soc Artif Organs 1956;2:102-5

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## POSSIBLE USE FOR INTRA-DIALYZER TRANSFER OF RESULTS

- PERFORM PRELIMINARY *IN VITRO* STUDY TO OBTAIN P RATIO FOR DRUG & STANDARD COMPOUND FOR DIALYZER BEING USED IN DIALYSIS STUDY (RECORD Q & RBC/PLASMA).
- THIS RATIO CAN BE USED TO ESTIMATE DRUG CL<sub>D</sub> FOR OTHER DIALYZERS AND OTHER Q VALUES IF P OF STANDARD COMPOUND FOR THAT DIALYZER IS KNOWN.
- NEED TO SELECT APPROPRIATE STANDARD COMPOUND (? CREATININE).

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## RECOVERY CLEARANCE

### THE GOLD STANDARD

$$CL_D = \frac{C_D \cdot Vol_D}{A \cdot t}$$

$$CL_D = \frac{C_D \cdot Vol_D}{AUC_A}$$

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## A-V DIFFERENCE METHOD [FICK EQUATION]

$$CL = Q \left[ \frac{A - V}{A} \right]$$

$$E = \left[ \frac{A - V}{A} \right]$$

Q = DIALYZER BLOOD FLOW  
A = CONCENTRATION IN BLOOD COMING TO DIALYZER  
V = CONCENTRATION IN BLOOD LEAVING DIALYZER  
E = EXTRACTION RATIO

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## EXTRACTION RATIO

Renkin Equation :

$$E = 1 - e^{-P/Q}$$

Fick Equation :

$$E = \frac{A - V}{A}$$

In Each Case :

$$CL = Q \cdot E$$

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## TWO DIALYSIS MYTHS

- NEED TO USE BLOOD CONCENTRATIONS WHEN CALCULATING BLOOD CLEARANCE  
**BUT PLASMA CONCENTRATIONS PROPORTIONAL TO BLOOD CONCENTRATIONS, SO MAKES NO DIFFERENCE IN  $A/[A + V]$  RATIO**
- NEED TO USE PLASMA FLOW WHEN CALCULATING PLASMA CLEARANCE

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## PLASMA VS. BLOOD CLEARANCE

RECOVERY :  $CL_p = \frac{U \cdot V}{P}$

$CL_b = \frac{U \cdot V}{B}$

FICK :  $CL_p = Q_{PK} \left( \frac{A - V}{A} \right)$        $CL_b = Q_B \left( \frac{A - V}{A} \right)$

IF  $B > P$  :  $CL_p > CL_b$ , SO :  $Q_{PK} > Q_B > Q_p$

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## NAPA IN RBC IS DIALYZED

FLOW PARAMETER	MEAN VALUE mL/min
Q <sub>PK</sub>	223
Q <sub>MEAS</sub>	195 (p < 0.2)
Q <sub>EFF</sub> *	217 (p > 0.2)

$$* Q_{EFF} = [ (1 - Hct) + (RBC/P) (HCT) ] Q_{MEAS}$$

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## DIALYSIS SATURATION VS. RECOVERY CLEARANCE

DIALYSIS SATURATION ( $EC = C_d/C_p$ ):

$$CL_D = Q_d \frac{C_d}{C_p}$$

RECOVERY CLEARANCE :

$$CL_D = \frac{UV}{P\tau} = \frac{C_d V_d}{C_p \tau}$$

BUT :

$$Q_d = \frac{V_d}{\tau} \text{ SO EXPRESSIONS ARE EQUIVALENT}$$

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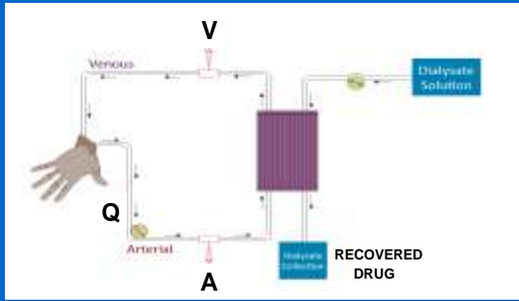
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## DATA SOURCES FOR FICK EQUATION




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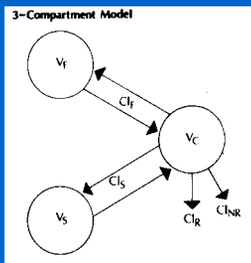
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## KINETIC MODEL USED TO ANALYZE HEMODIALYSIS DATA\*



\* From Stec GP, et al. Clin Pharmacol Ther 1979;26:618-28.

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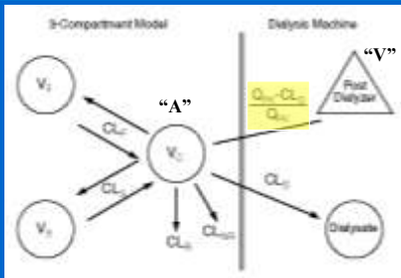
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\* From Stec GP, et al. Clin Pharmacol Ther 1979;26:618-28.

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## FICK CLEARANCE EQUATION

$$CL = Q \left[ \frac{A - V}{A} \right]$$

$$CLA = QA - QV$$

$$QV = QA - CLA$$

$$V = \left[ \frac{Q - CL}{Q} \right] A$$

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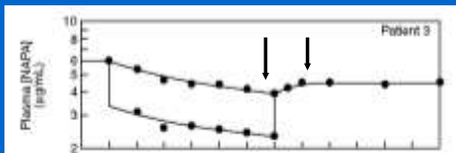
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## TWO PROBLEMS WITH FIXED-PARAMETER MODEL\*



1. DURING DIALYSIS: [A] AND [V] DROP MORE THAN EXPECTED FROM DRUG RECOVERY
2. AFTER DIALYSIS: CONCENTRATION REBOUND IS LESS THAN EXPECTED

\* From Stec GP, et al. Clin Pharmacol Ther 1979;26:618-28.

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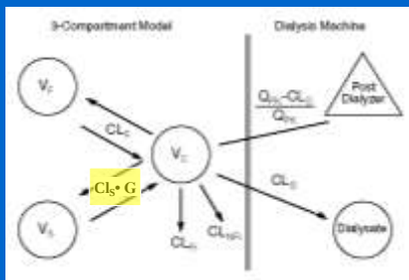
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\* From Stec GP, et al. Clin Pharmacol Ther 1979;26:618-28.

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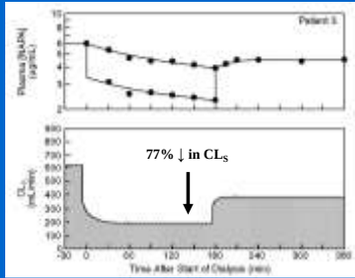
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## REDUCTION IN $CL_S$ DURING AND AFTER HEMODIALYSIS\*



\* From Stec GP, et al. Clin Pharmacol Ther 1979;26:618-28.

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## CONDUCT OF PK STUDIES IN HEMODIALYSIS PATIENTS

Chapter 6 – Principles of Clinical Pharmacology

Atkinson AJ Jr, Umans JG: Pharmacokinetic Studies in Hemodialysis Patients. Clin Pharmacol Ther 2009;86:548-52.

CDER, FDA: Draft Guidance for Industry – Pharmacokinetics in Patients with Impaired Renal Function – Study Design, Data Analysis, and Impact on Dosing and Labeling. <http://www.fda.gov/downloads/Drugs/GuidanceComplianceRegulatoryInformation/Guidances/UCM204959.pdf>

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### CASE HISTORY

A 67 year-old woman became lethargic and confused and developed hypotension, renal insufficiency, junctional tachycardia and intraventricular conduction delay after ingesting an estimated 7gm of procainamide (PA). Plasma PA and NAPA concentrations were 57 µg/mL and 55 µg/mL, respectively.

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### CASE HISTORY (cont.)

Hemodialysis was performed for 4 hr. By the end of the second hour BP was maintained in the range of 110/80 mm Hg without vasopressor therapy. At the end of dialysis, the patient was alert and oriented although only 340 mg of PA and 470 mg of NAPA had been removed by this procedure.

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### DIALYSIS CASE HISTORY (cont.)

Fifteen hours after dialysis, PA and NAPA levels were 9.2 µg/mL and 33 µg/mL, respectively. The patient had returned to normal sinus rhythm with QRS = 0.12 sec.

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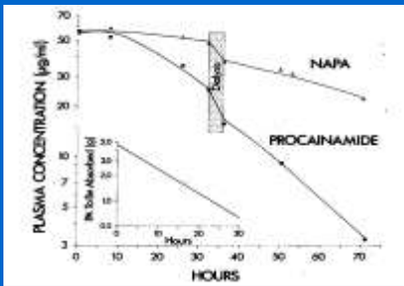
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## KINETIC ANALYSIS OF HEMODIALYSIS FOR PROCAINAMIDE TOXICITY\*



\* From: Atkinson AJ Jr, et al. Clin Pharmacol Ther 1976;20:585-92.

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## CRITERION FOR DIALYSIS EFFICACY\*

$$CL_{EC} > 30\% [CL_R + CL_{NR}]$$

\* Levy G. Am J Med 1977;62:461-5.

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## WAS DIALYSIS EFFICACIOUS?

- **DIALYSIS INCREASED DRUG CLEARANCE**  
 PA – TWO FOLD  
 NAPA – 3.8 FOLD
  - **BUT 4 hr OF DIALYSIS REMOVED < 1 gm of 7 gm DOSE**  
 340 mg PA  
 470 mg NAPA
  - **HOWEVER, BLOOD LEVELS FELL SUBSTANTIALLY**  
 PA: 25.7 µg/mL → 15.5 µg/mL  
 NAPA: 47.0 µg/mL → 35.5 µg/mL
- AND PATIENT'S CONDITION STABILIZED**

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## PA & NAPA KINETICS IN TOXIC PATIENT

	NORMAL		PATIENT	
	PA	NAPA	PA	NAPA
$t_{1/2}$ (hr)	2.5	6.2	10.5	35.9
$CL_E$ (mL/min)	590	233	66.8	16.1
$CL_D$ (mL/min)			68.3	45.8
$V_{d\beta}$ (L/kg)	1.80	1.76	0.76	0.63

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## ESTIMATION OF $V_d$

**Question: Why was distribution volume estimate so much lower in patient than in normal subjects?**

USUAL  $V_d$  ESTIMATE :

$$V_d = \frac{\text{DOSE GIVEN}}{\Delta \text{ CONCENTRATION}}$$

DIALYSIS  $V_d$  ESTIMATE :

$$V_d = \frac{\text{DRUG REMOVED}}{\Delta \text{ CONCENTRATION}}$$

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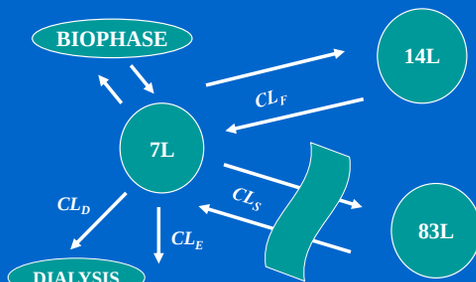
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## SEQUESTRATION OF DRUG IN SOMATIC TISSUES




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### EFFICACY OF EXTRACORPOREAL TREATMENT OF DRUG TOXICITY

- TOTAL EXTENT OF DRUG REMOVAL MAY BE COMPROMIZED BY ↓ CL<sub>s</sub>.
- ↓ CL<sub>s</sub> FROM SOMATIC TISSUES CAN ACCELERATE ↓ IN DRUG CONCENTRATION TO WHICH VITAL ORGANS (CNS, HEART) ARE EXPOSED AND RESULT IN A BENEFICIAL CLINICAL RESPONSE > EXTENT OF DRUG REMOVAL.
- ↓ CL<sub>s</sub> FROM SOMATIC TISSUES ALSO ATTENUATES POST-DIALYSIS REBOUND.

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### WHY DOES CL<sub>s</sub> ↓ DURING DIALYSIS ?

$$CL = Q(1 - e^{-P \cdot S/Q})$$

#### POSSIBILITIES:

- CAPILLARY BLOOD FLOW (Q) DECREASES
- CAPILLARY P · S PRODUCT DECREASES
- BOTH DECREASE

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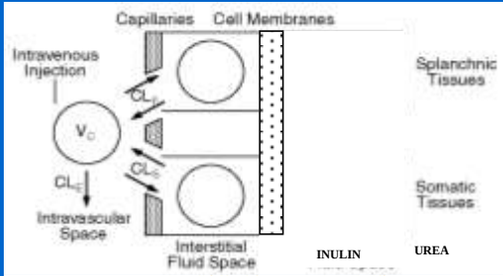
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**MULTICOMPARTMENTAL MODEL OF INULIN AND UREA KINETICS\***



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

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**BASIS FOR KINETIC HETEROGENEITY OF INTERSTITIAL FLUID SPACE**

EFFECTIVE PORE SIZE	CAPILLARY STRUCTURE	PRIMARY LOCATION
LARGE	FENESTRATED	SPLANCHNIC BED
SMALL	CONTINUOUS	SOMATIC TISSUES

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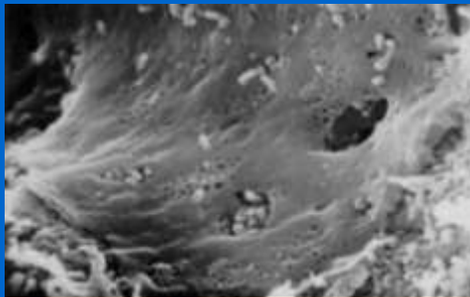
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**ENDOTHELIAL FENESTRAE IN HEPATIC SINUSOIDS**




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**INTERENDOTHELIAL CELL JUNCTION IN CONTINUOUS CAPILLARY**




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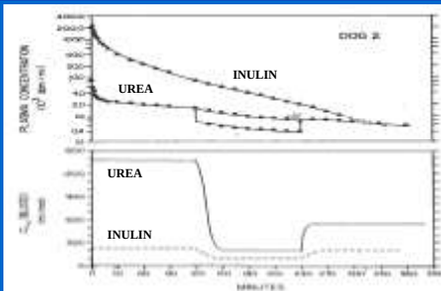
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**UREA (●) AND INULIN (▲) KINETICS DURING AND AFTER HEMODIALYSIS\***



\* From Bowsher DJ, et al. J Lab Clin Med 1985;105:489-97.

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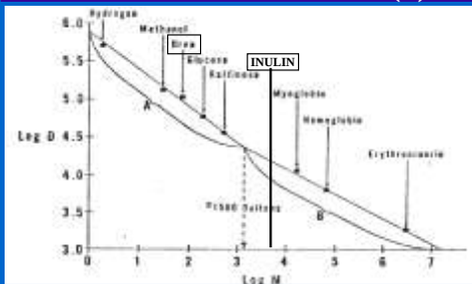
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**EFFECT OF MOLECULAR WEIGHT (M) ON SOLUTE DIFFUSIVITY (D)\***



\* From Henderson LW: In: Brenner BM, Rector FC Jr. The Kidney. 1976, p. 1643-71.

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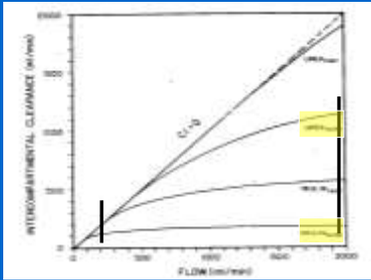
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## RELATIONSHIP BETWEEN BLOOD FLOW (Q) AND $CL_I$ \*



\* From Bowsheer DJ, et al. J Lab Clin Med 1985;105:489-97.

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## UREA AND INULIN KINETICS DURING AND AFTER HEMODIALYSIS

PARAMETER	BEFORE	DURING	AFTER
<b>BLOOD FLOW</b>			
$Q_S$ (mL/min)	1991	199	405
$Q_F$ (mL/min)	2332	2591*	2965*
C.O. (mL/min)	4399	2790	3370
<b>PS</b>			
INULIN (mL/min)	186	169	238
UREA (mL/min)	1649	1541	2164

\* ESTIMATED AS C.O. -  $Q_S$

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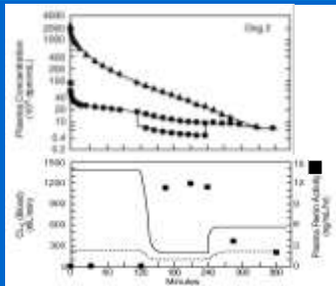
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## RENIN-ANGIOTENSIN SYSTEM ACTIVATION DURING AND AFTER HEMODIALYSIS\*



\* From Bowsheer DJ, et al. J Lab Clin Med 1985;105:489-97.

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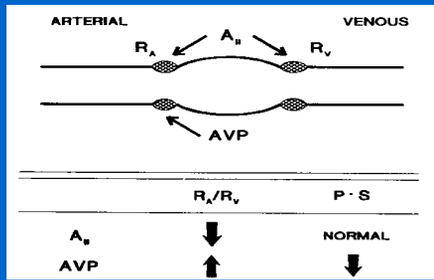
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## DIFFERENT MICROCIRCULATORY ACTIONS OF ANGIOTENSIN II AND AVP\*



\* From Atkinson AJ Jr: The Pharmacologist 1989;31:229-34.

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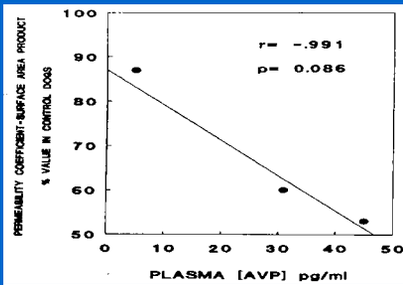
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## EFFECT OF ARGININE VASOPRESSIN (AVP) ON $P \cdot S$ \*



\* From Atkinson AJ Jr: The Pharmacologist 1989;31:229-34.

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## HEMODIALYSIS-ASSOCIATED SKELETAL MUSCLE CRAMPS

- COMPLICATE MORE THAN 20% OF HEMODIALYSIS SESSIONS
- PATHOGENESIS UNCLEAR
- SYMPTOMATIC THERAPY: NaCl, MANNITOL
- PREVENTIVE THERAPY: NaCl INFUSION
- OCCUR MORE FREQUENTLY IN SOME PATIENTS THAN OTHERS

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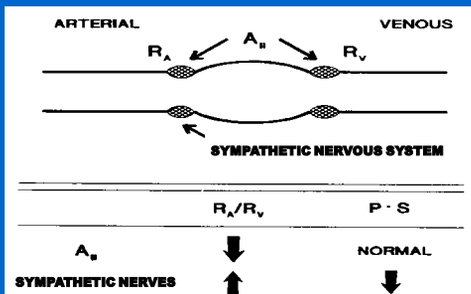
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## ACTIONS OF ANGIOTENSIN II & SYMPATHETIC NERVOUS SYSTEM




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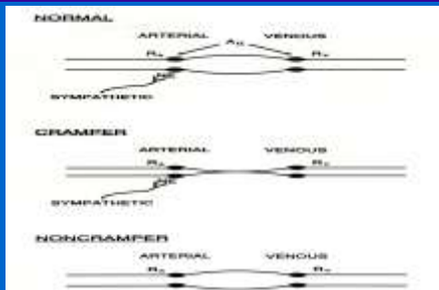
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## ONLY SOME PATIENTS HAVE DIALYSIS-ASSOCIATED SKELETAL MUSCLE CRAMPS\*



\* Sidhom OA, et al. Clin Pharmacol Ther 1994;56:445-51

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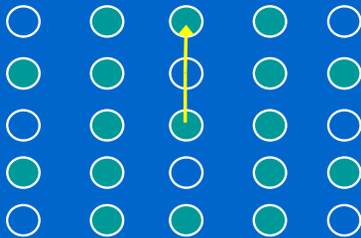
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### CAPILLARY DERECRUITMENT (OPEN (O) & CLOSED (●) CAPILLARIES)



8 OPEN CAPILLARIES IN MUSCLE CROSS SECTION

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### PATHOGENESIS OF DIALYSIS-ASSOCIATED SKELETAL MUSCLE CRAMPS



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### CONCLUDING THOUGHT

ALTHOUGH NON-COMPARTMENTAL ANALYSIS OF PK DATA IS CURRENTLY IN VOGUE, IT IS UNABLE TO PROVIDE INSIGHT INTO SOME IMPORTANT PHENOMENA:

- IMPACT OF DIALYSIS-ASSOCIATED HEMODYNAMIC CHANGES ( $\downarrow$   $CL_s$ )
- IMPACT OF  $\downarrow$  SPLANCHNIC BLOOD FLOW ( $\downarrow$   $CL_p$ ) ON BIOAVAILABILITY

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