

GUYTON AND HALL *Textbook of*
Medical Physiology

TWELFTH EDITION



Chapter 29:

Renal Regulation of Potassium, Calcium, Phosphate,
and Magnesium; Integration of Renal Mechanisms for
Control of Blood Volume and Extracellular Fluid Volume



Normal potassium intake, distribution, and output from the body.

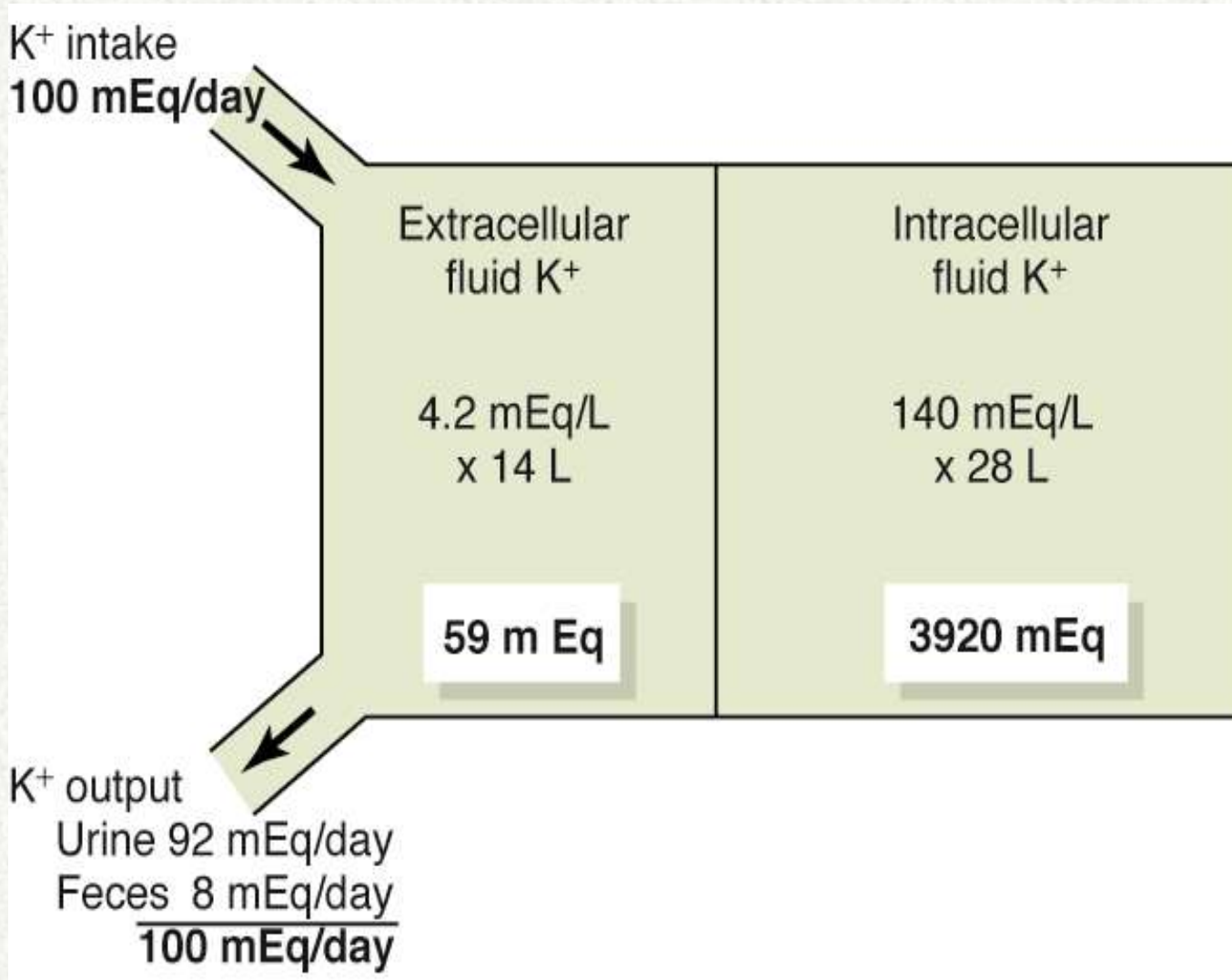


Figure 29-1



Effects of severe hyperkalemia

- Partial depolarization of cell membranes
- Cardiac toxicity
 - ventricular fibrillation or asystole

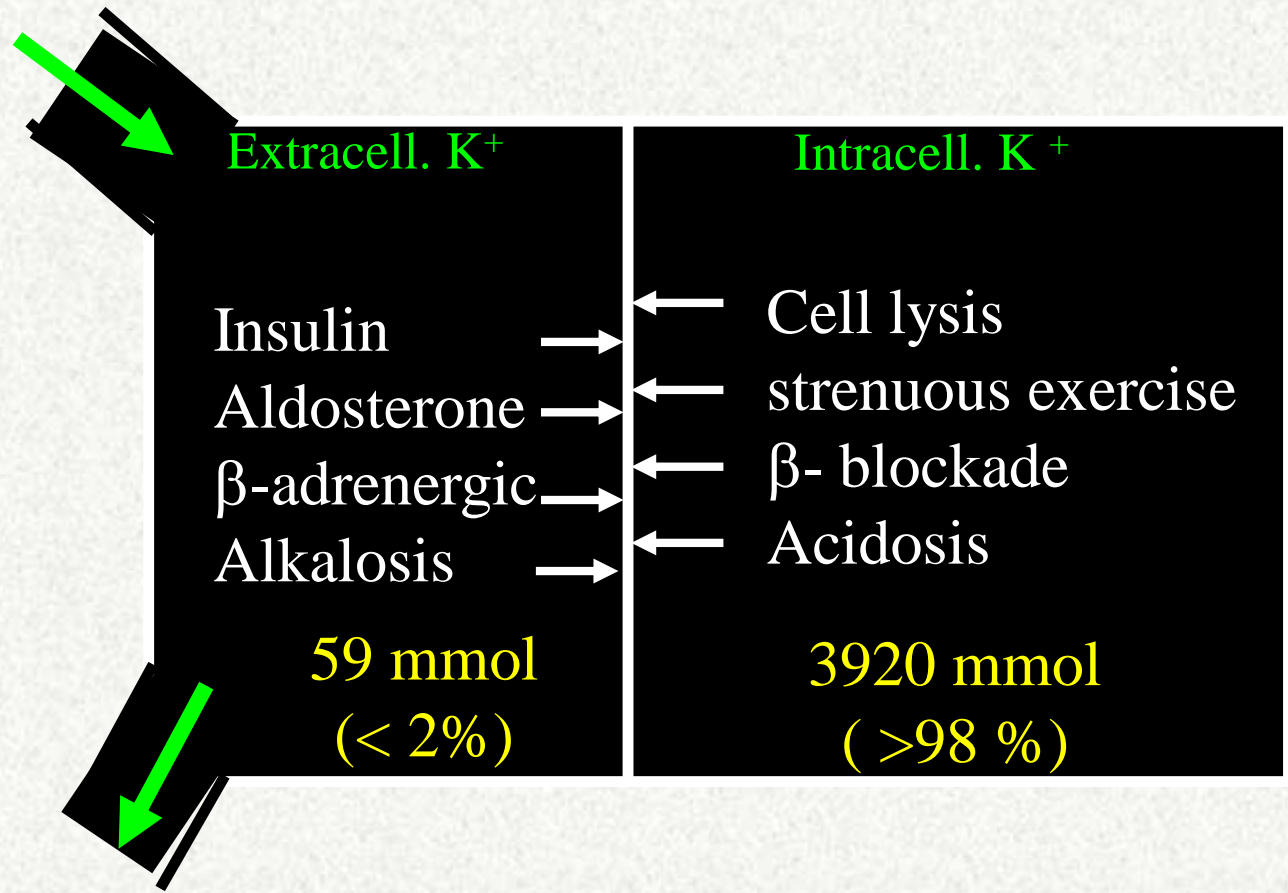
Effects of severe hypokalemia

- Hyperpolarization of cell membranes
- Fatigue, muscle weakness
- hypoventilation
- delayed ventricular repolarization



Potassium Regulation: Internal and External

K^+ intake
100 mEq/d



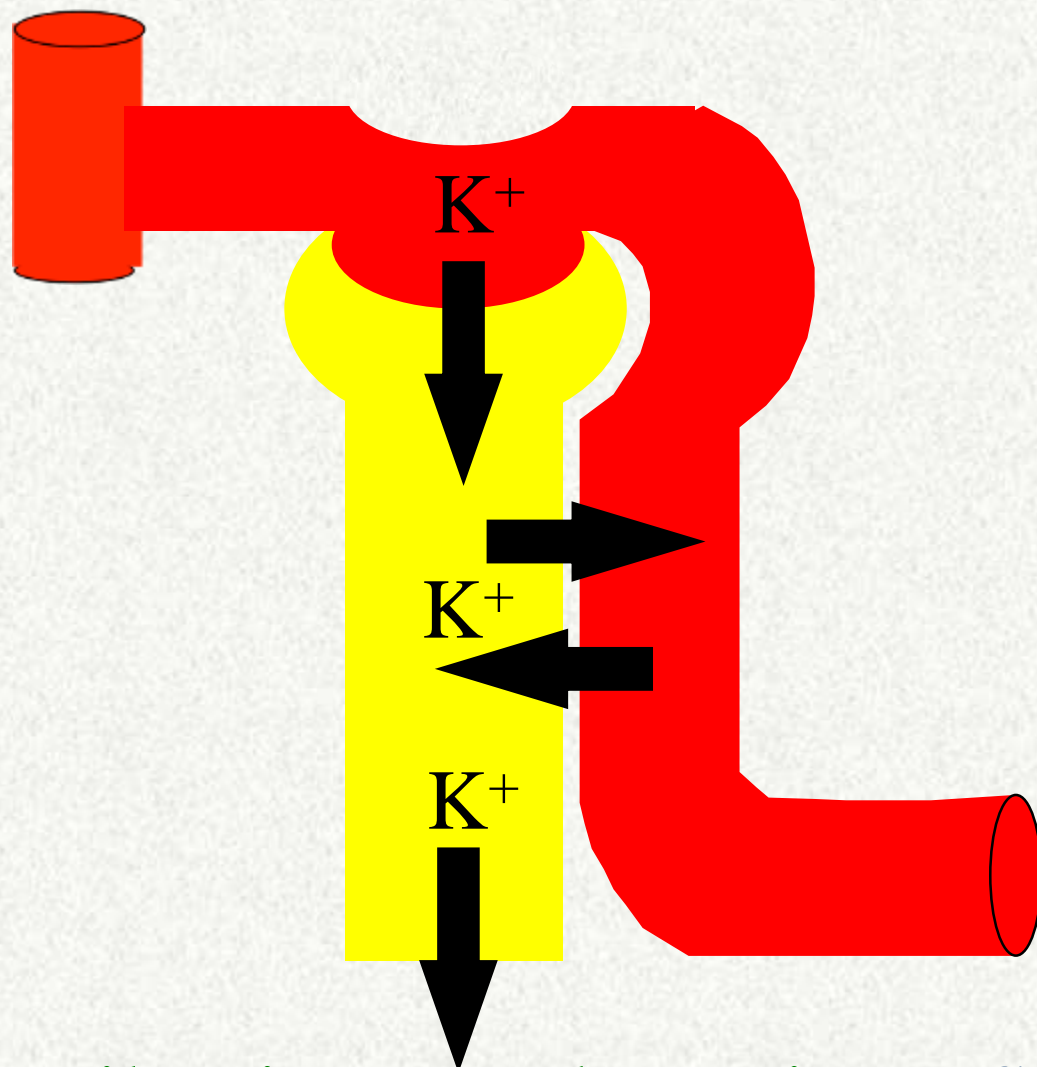
59 mmol
(< 2%)

3920 mmol
(> 98 %)

K^+ output
Total = 100 mEq/d



Control of Potassium Excretion



$$\text{Excretion} = \text{Filtration} - \text{Reabsorption} + \text{Secretion}$$



Renal tubular sites of potassium reabsorption and secretion.

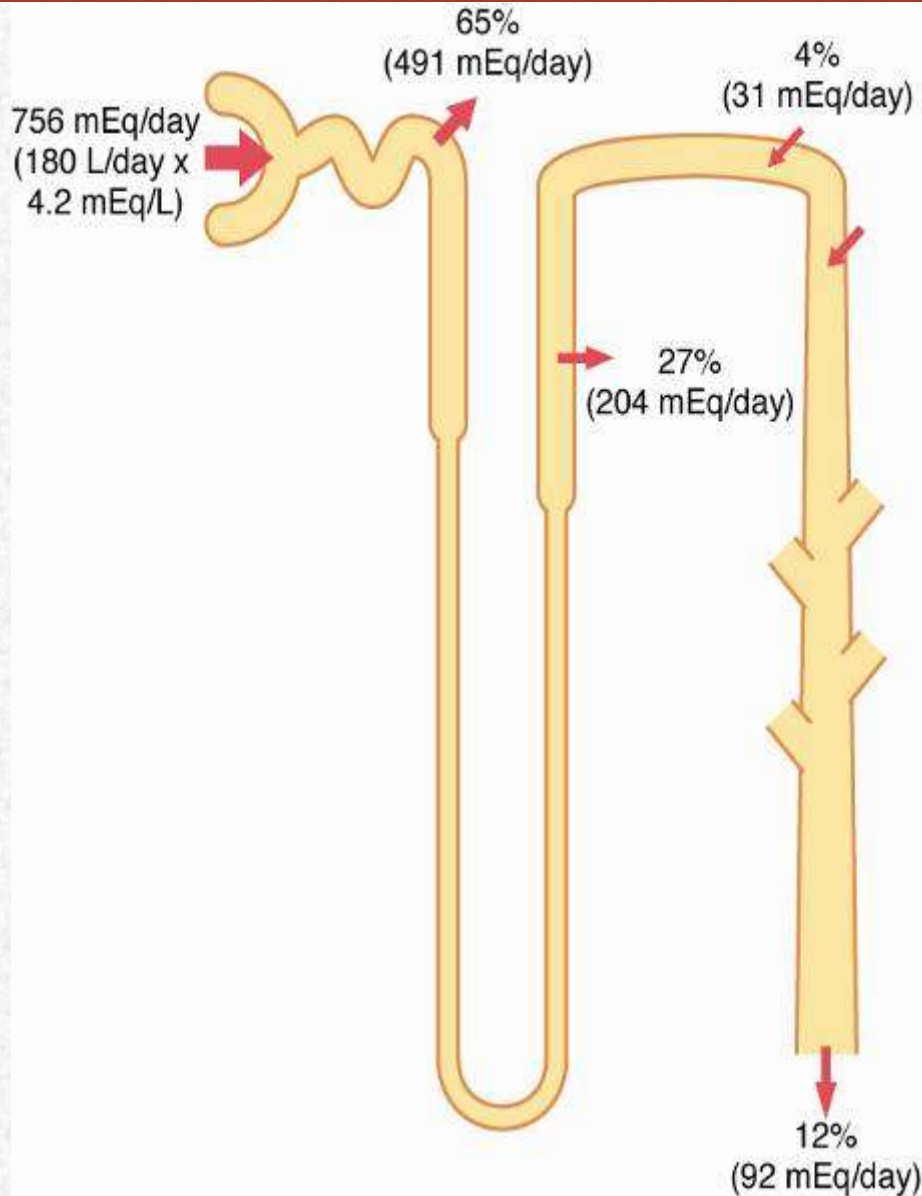
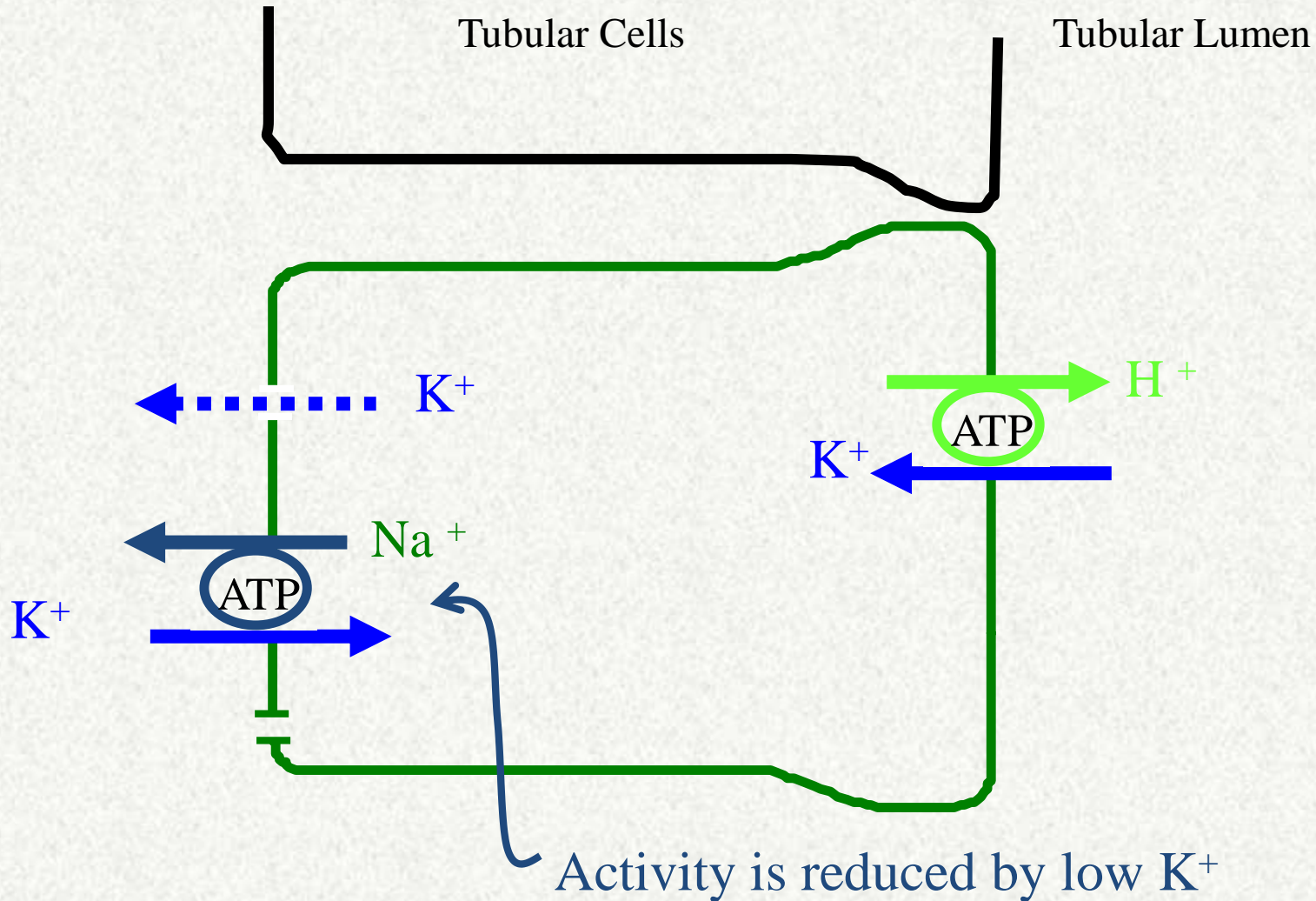


Figure 29-2



Late Distal and Cortical Collecting Tubules Intercalated Cells – Reabsorb K^+





Potassium Secretion by Principal Cells

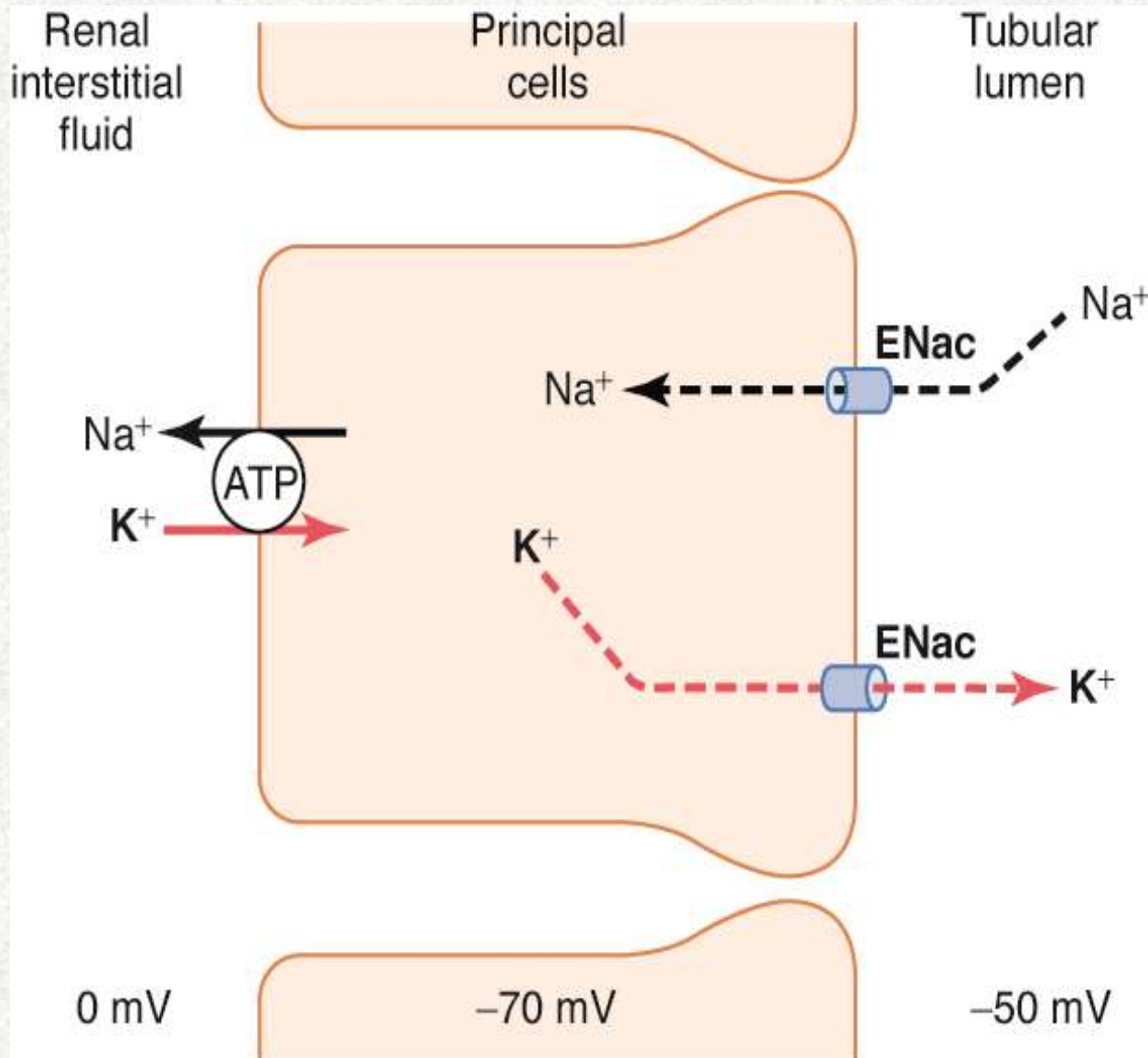


Figure 29-3



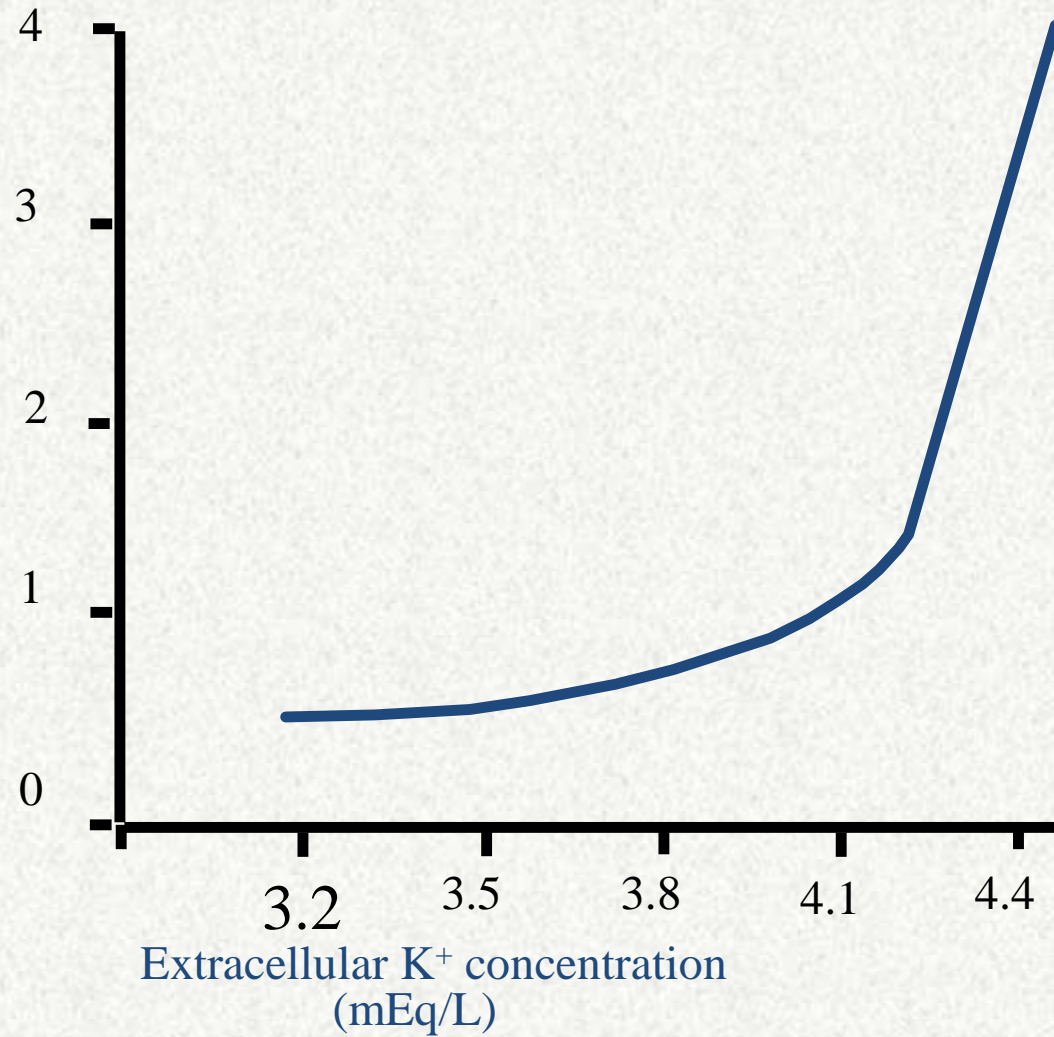
Control of Cortical Collecting Tubule (Principal Cells) K^+ Secretion

- Extracellular K^+ concentration : increases K^+ secretion
- Aldosterone : increases K^+ secretion
- Sodium (volume) delivery : increases K^+ secretion
- Acid - base status:
 - acidosis : decreases K^+ secretion
 - alkalosis : increases K^+ secretion



Effect of Extracellular K^+ on Excretion of K^+

Tubular
 K^+
Secretion
(x normal)

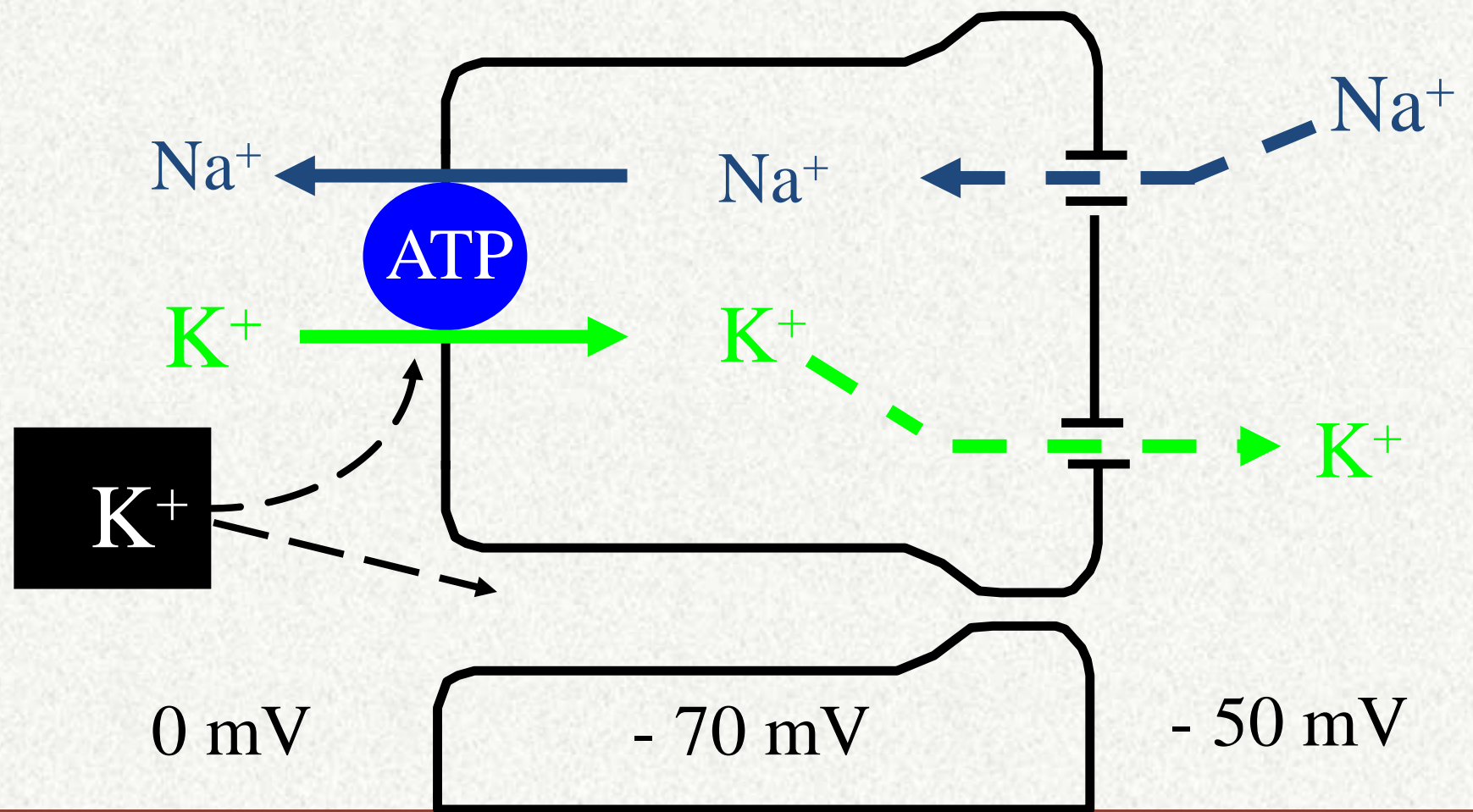




Interstitial Fluid

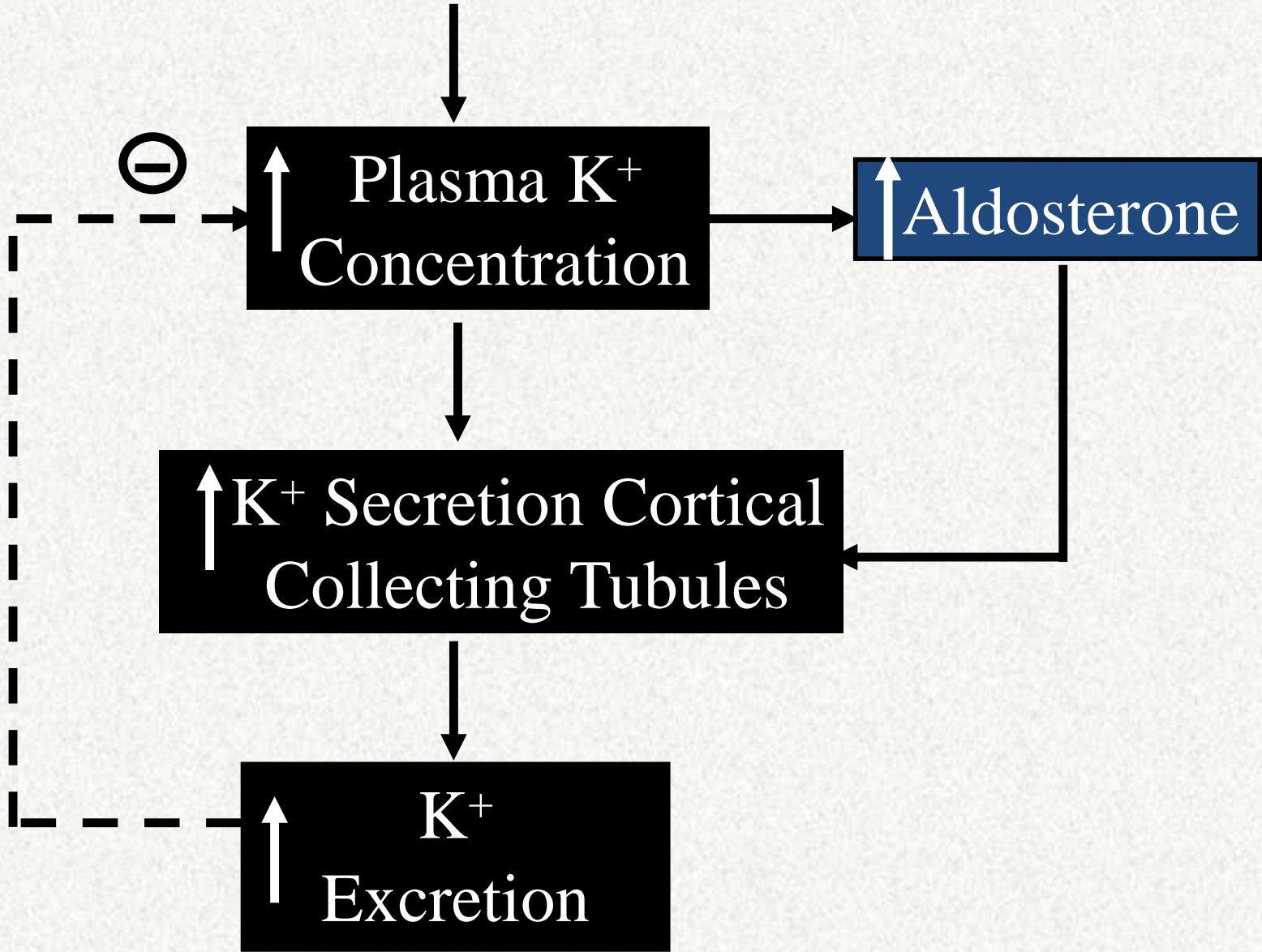
Principal Cells

Tubular Lumen





↑ K^+ Intake





Increased serum K^+ stimulates aldosterone secretion

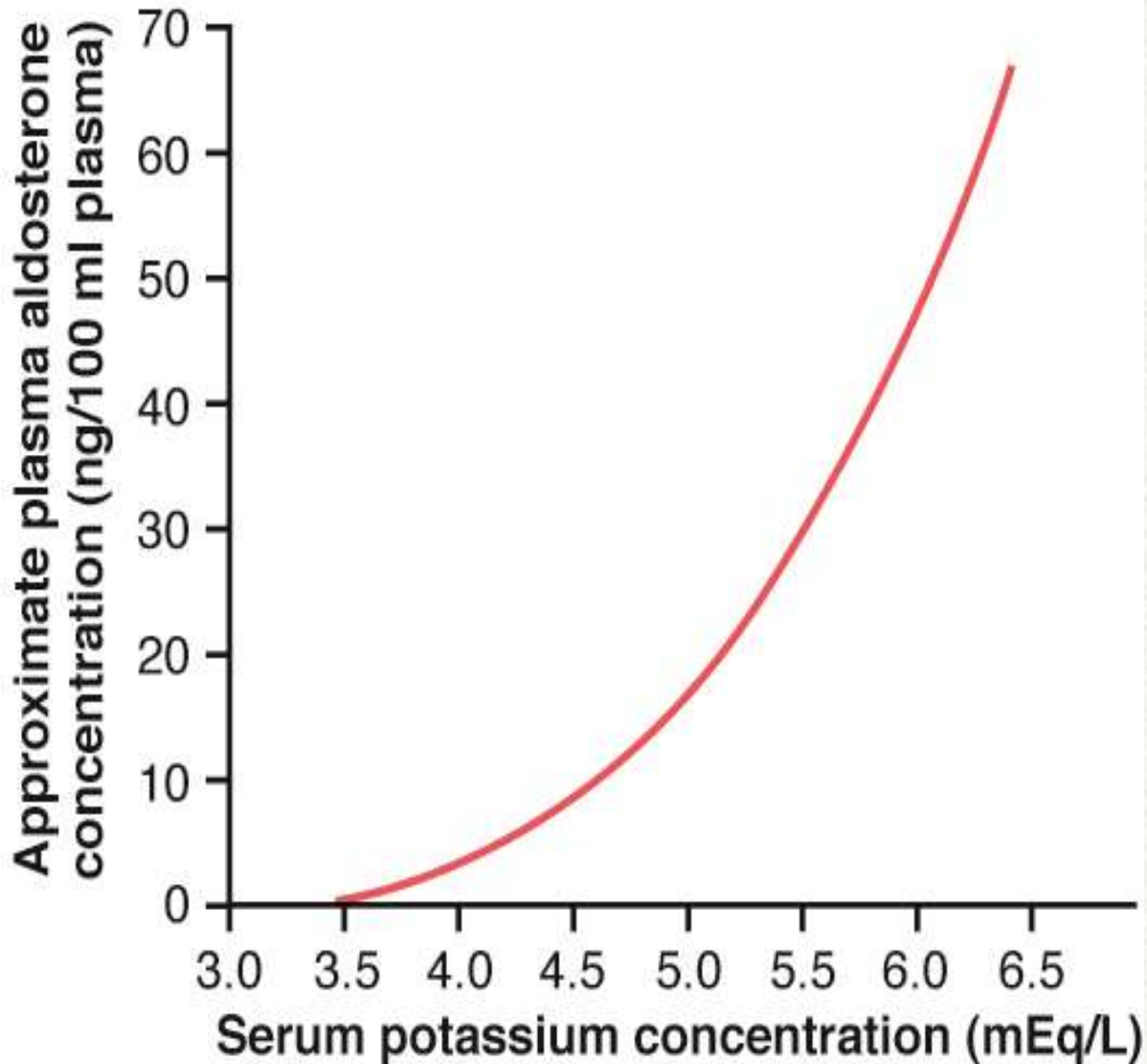
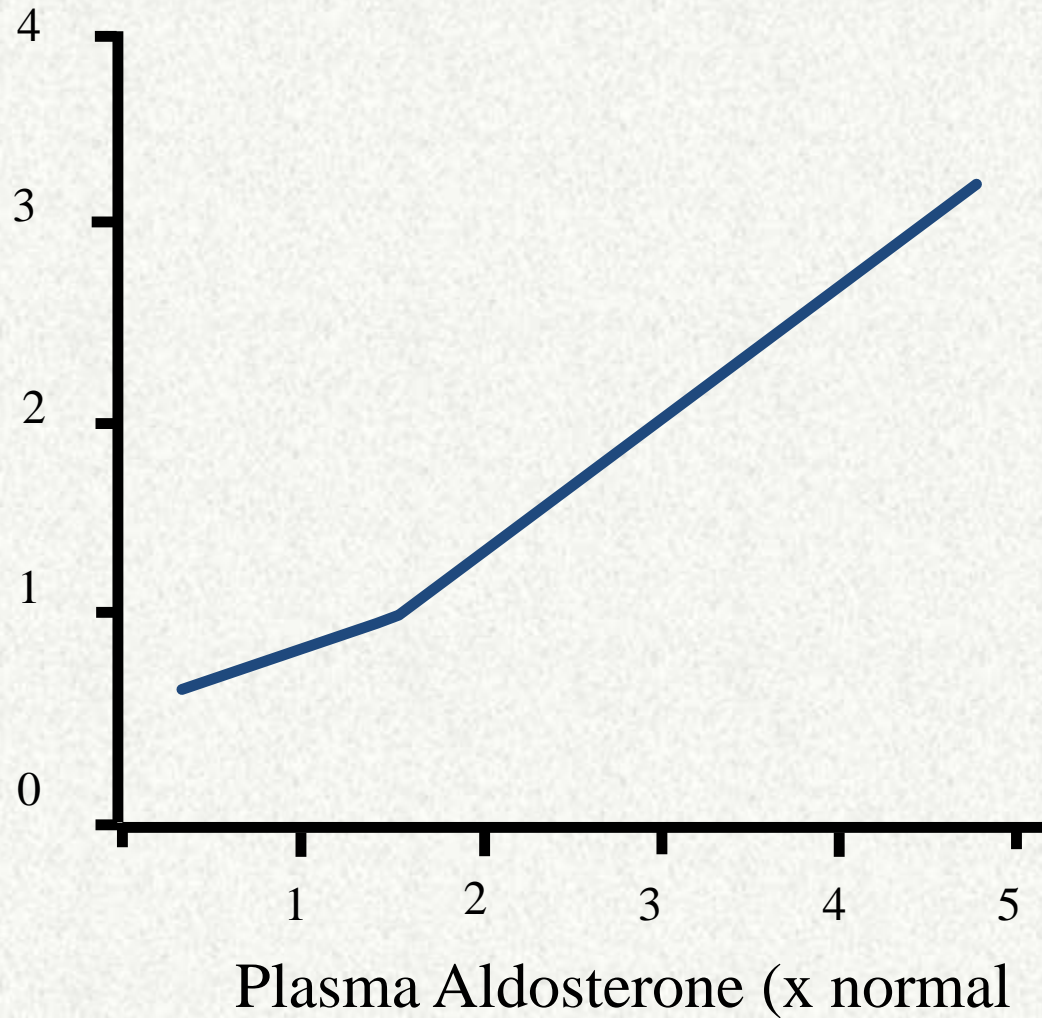


Figure 29-5



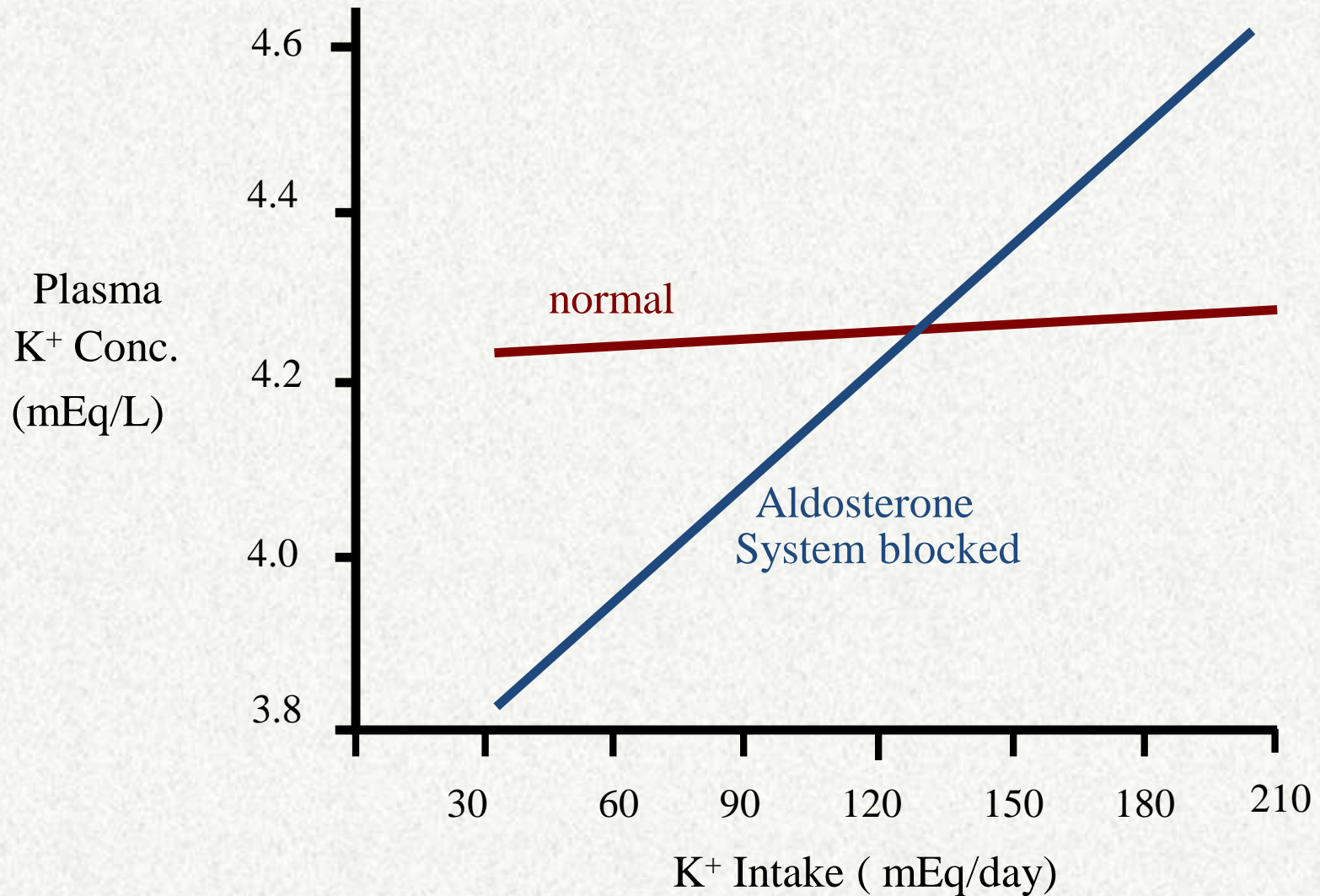
Effect of Aldosterone on K^+ Excretion

Tubular
 K^+
Secretion
(x normal)





Effect of Changes in K^+ Intake on Plasma K^+ After Blocking Aldosterone System





Effect of collecting tubule flow rate on K^+ secretion

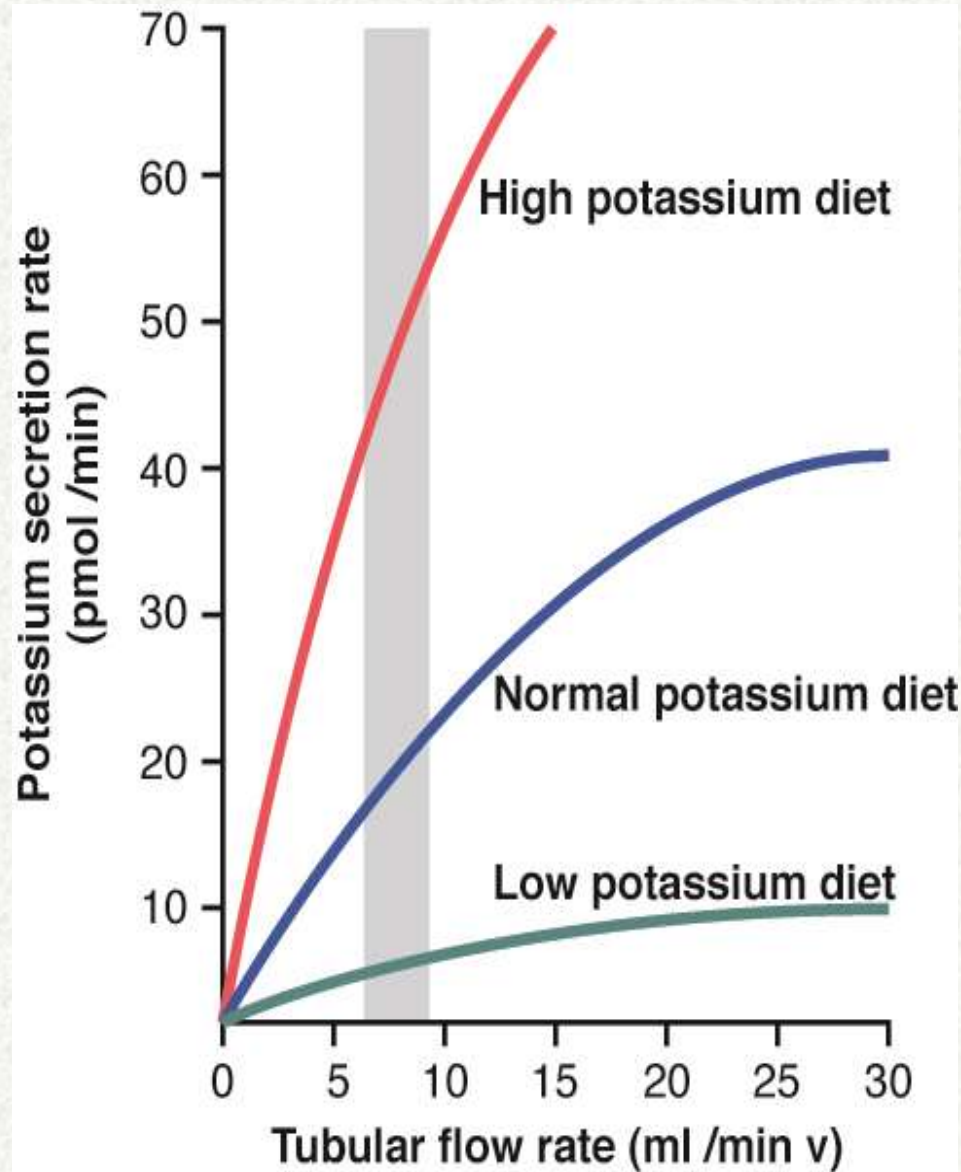


Figure 29-9



Diuretics that ↓ Prox. or Loop Na⁺ Reabsorption

↓
Water Reabsorption

↑
Volume Delivery to Cort. Collect. Tub.

↑
Cell : Lumen Gradient for K⁺ Diffusion

↑
K⁺ Secretion

K⁺ Depletion

↓
K⁺ Reabsorption



↑ Na⁺ Intake

↓ Aldosterone

↑ GFR

↓ Proximal Tubular Na⁺ Reabsorption

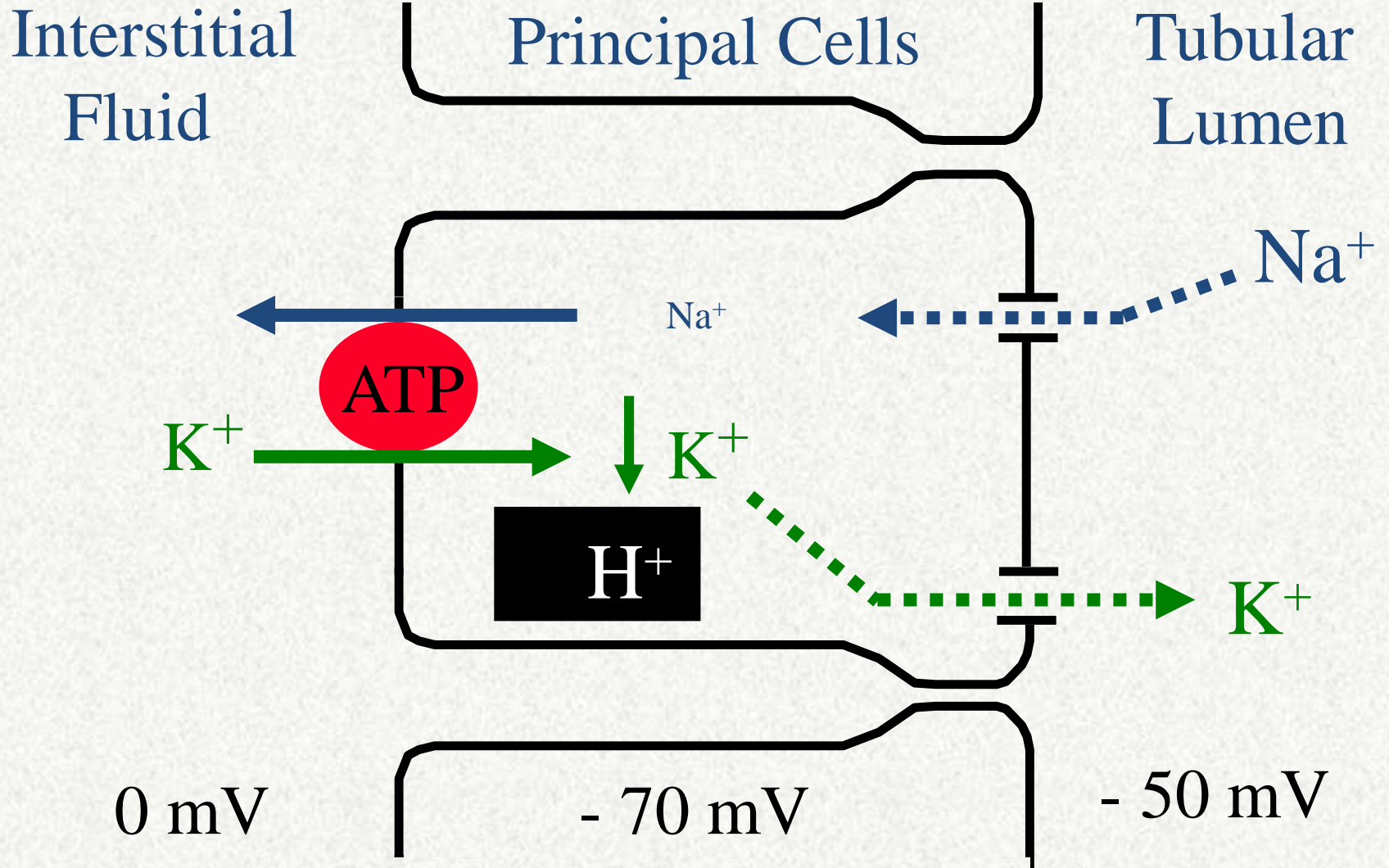
↑ K⁺ Secretion
Cort. Collect. Ducts

↑ Distal Tubular Flow Rate

Unchanged
K⁺ Excretion

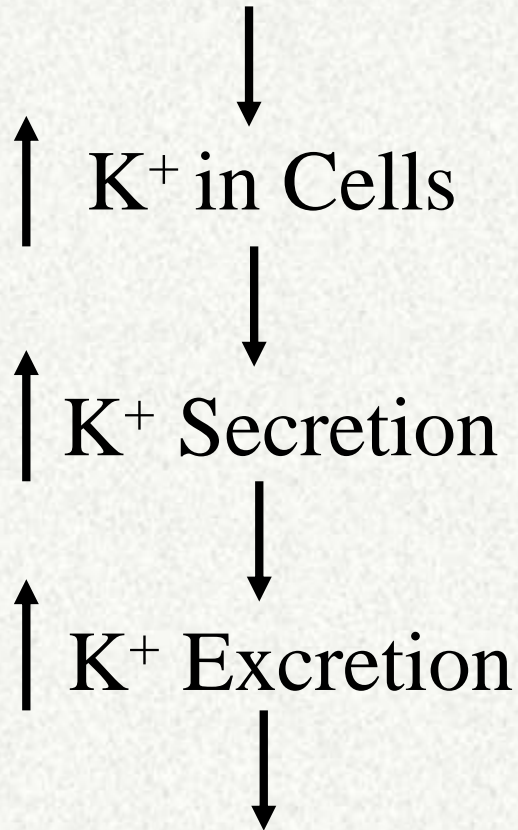


Acidosis Decreases Cell K^+





Alkalosis



K⁺ Depletion



Causes of Hyperkalemia

- Renal failure
- Decreased distal nephron flow (heart failure, severe volume depletion, NSAID, etc)
- Decreased aldosterone or decreased effect of aldosterone
 - adrenal insufficiency
 - K^+ sparing diuretics (spironolactone, eplerenone)
- Metabolic acidosis (hyperkalemia is mild)
- Diabetes (kidney disease, acidosis, ↓ insulin)



Causes of Hypokalemia

- Very low intake of K^+
- GI loss of K^+ - diarrhea
- Metabolic alkalosis
- Excess insulin
- Increased distal tubular flow /
 - salt wasting nephropathies
 - osmotic diuretics
 - loop diuretics
- Excess aldosterone or other mineralocorticoids



Compensatory responses to decreased plasma ionized calcium

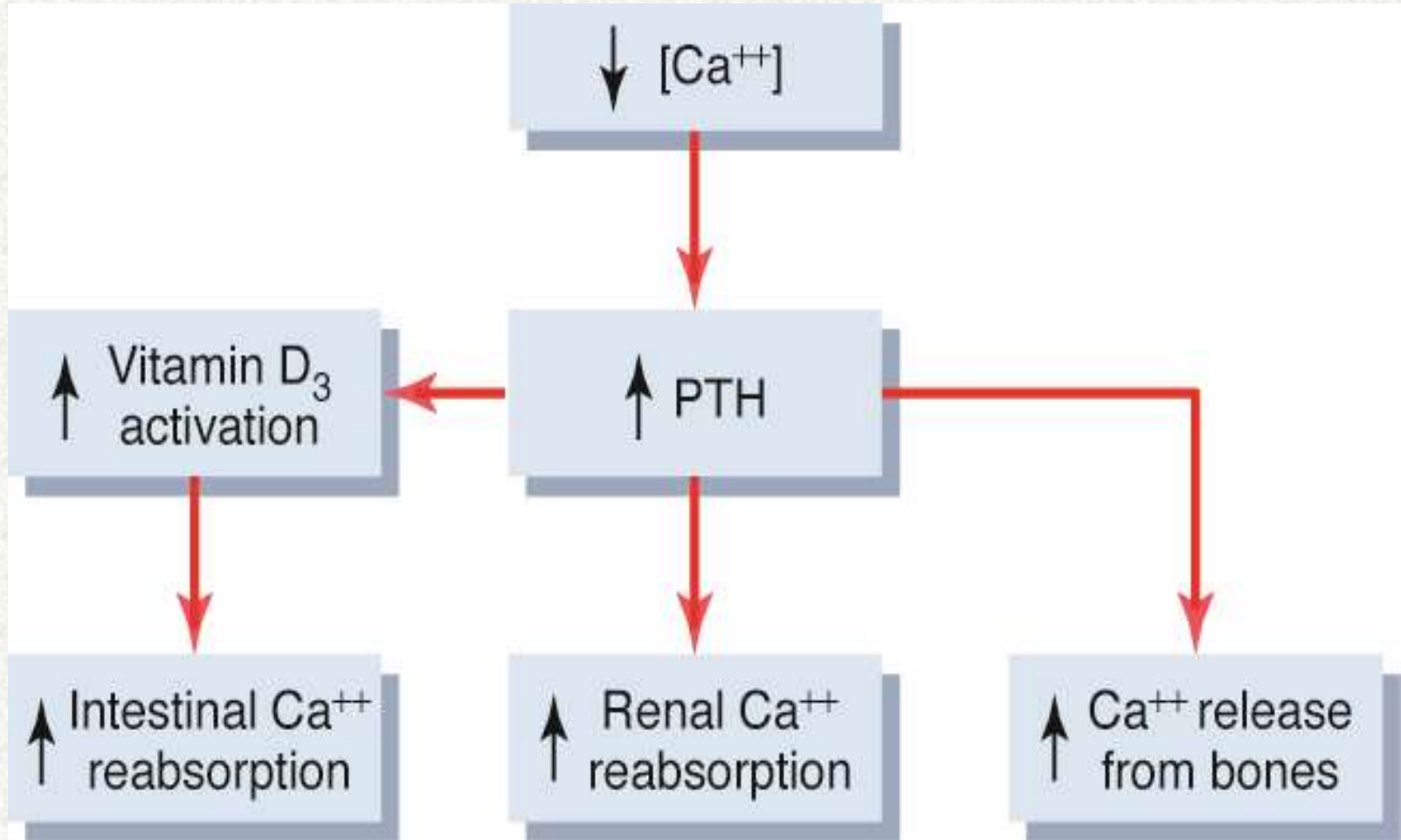


Figure 29-11



Proximal tubular calcium reabsorption

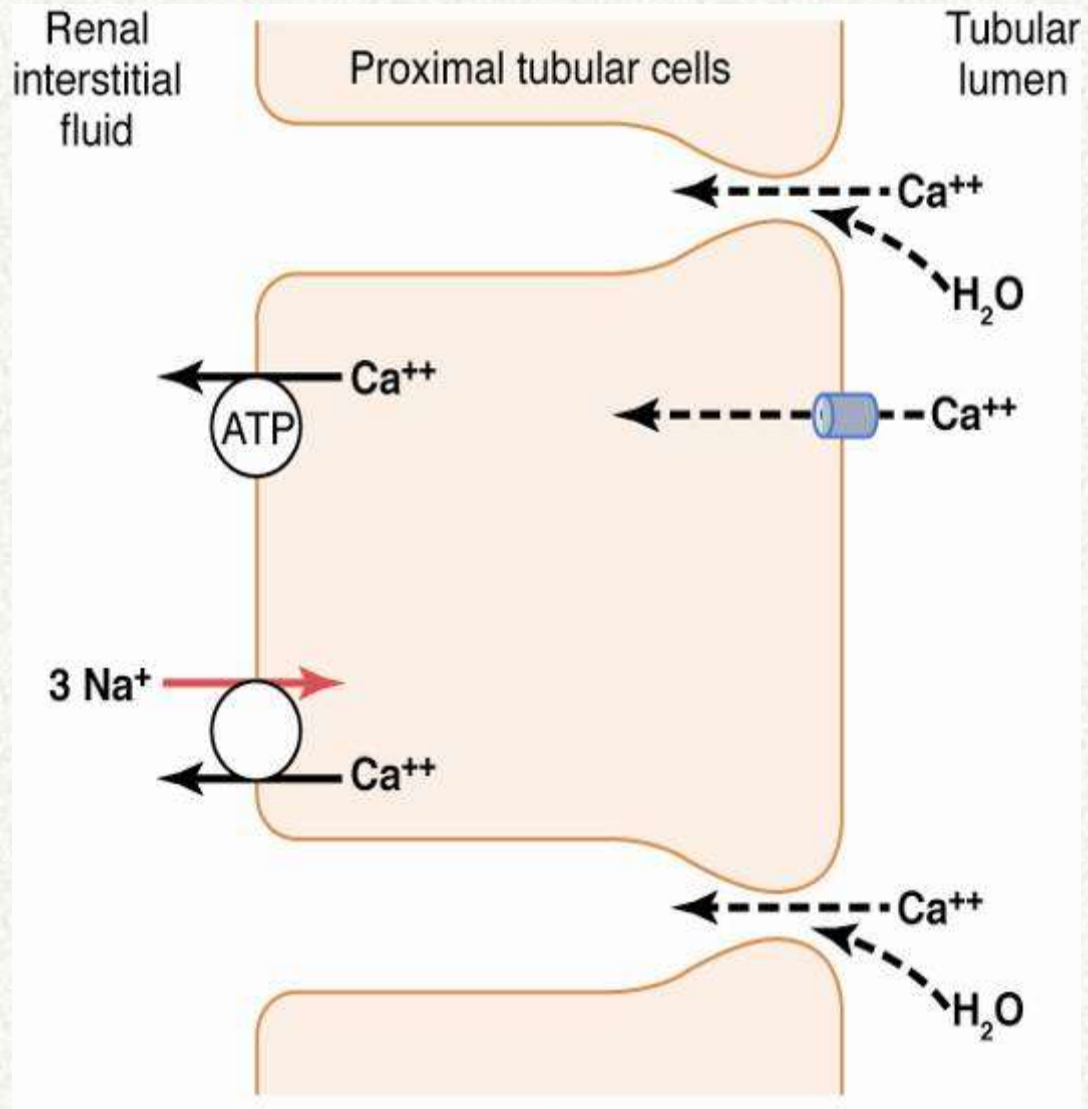


Figure 29-12



Integration of Renal Mechanisms for Regulation of Body Fluids

$\text{Excretion} = \text{Filtration} - \text{Reabsorption} + \text{Secretion}$

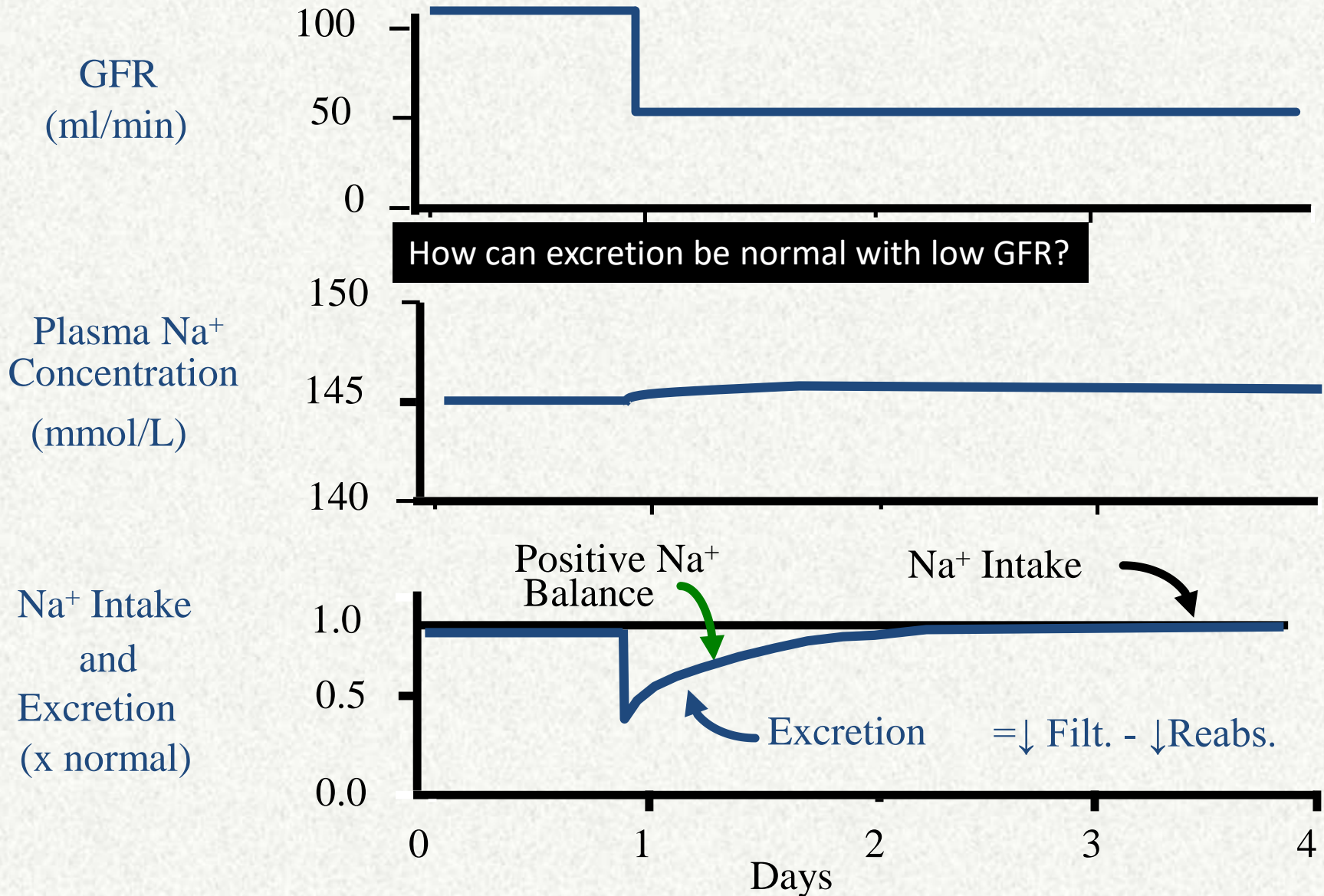
If there is a steady - state :

$\text{Fluid Excretion} = \text{Fluid Intake}$

$\text{Electrolyte Excretion} = \text{Electrolyte intake}$

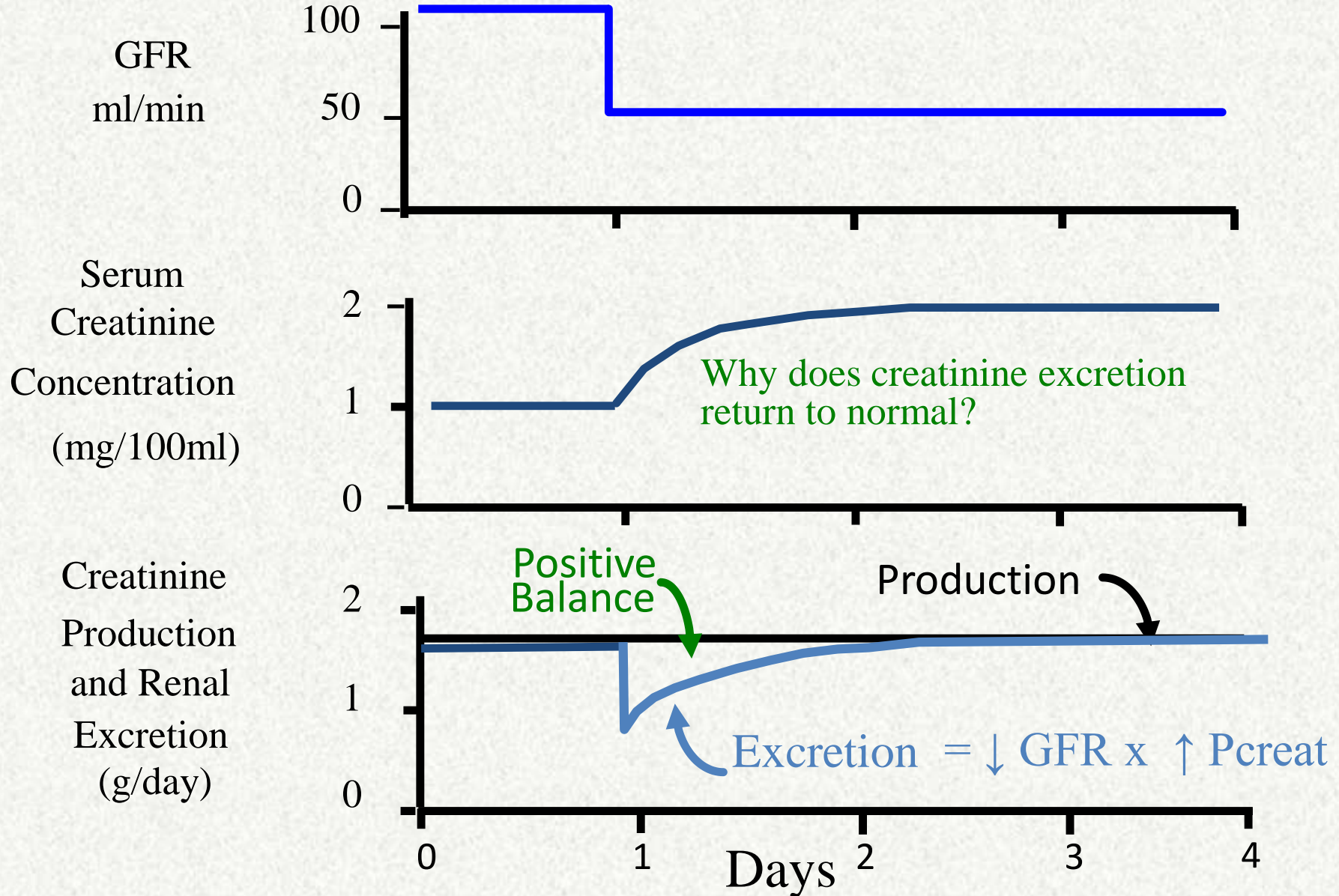


Effect of Decreased GFR on Sodium





Effect of Decreased GFR on Creatinine





Plasma concentrations of solutes in chronic renal failure

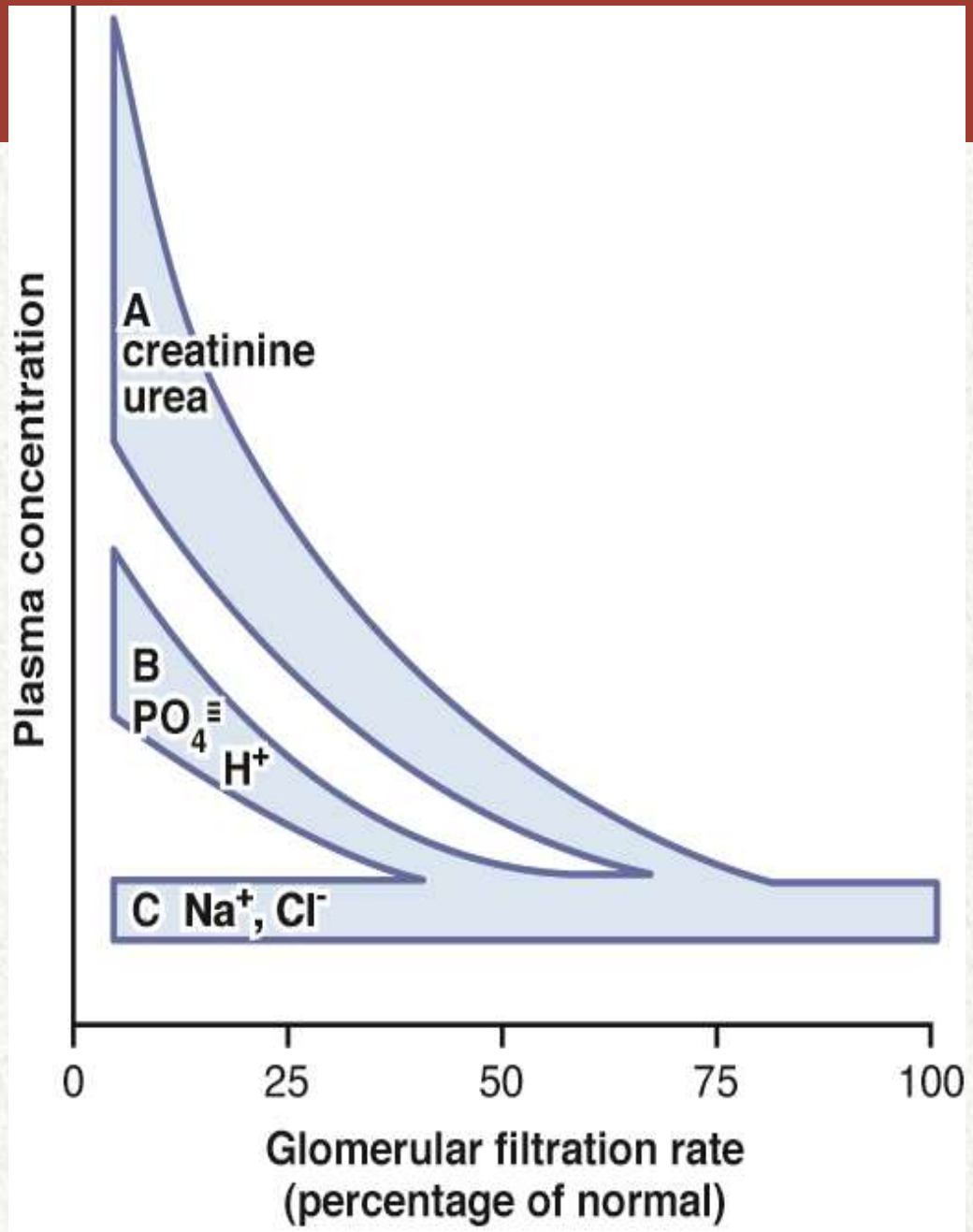


Figure 31-5



Hierarchy of Responses to Disturbances of Body Fluid Regulation

1. Local renal mechanisms

- changes in GFR
- changes in tubular reabsorption
- changes in tubular secretion

2. Systemic mechanisms (which can affect the whole body)

- changes in hormones
- changes in sympathetic activity
- changes in blood pressure
- changes in blood composition



↓ GFR

Proximal Delivery.

Loop NaCl Delivery

Macula Densa NaCl Delivery

Distal NaCl Delivery

Urine Excretion
Na⁺ *Minimized*

Reabsorption

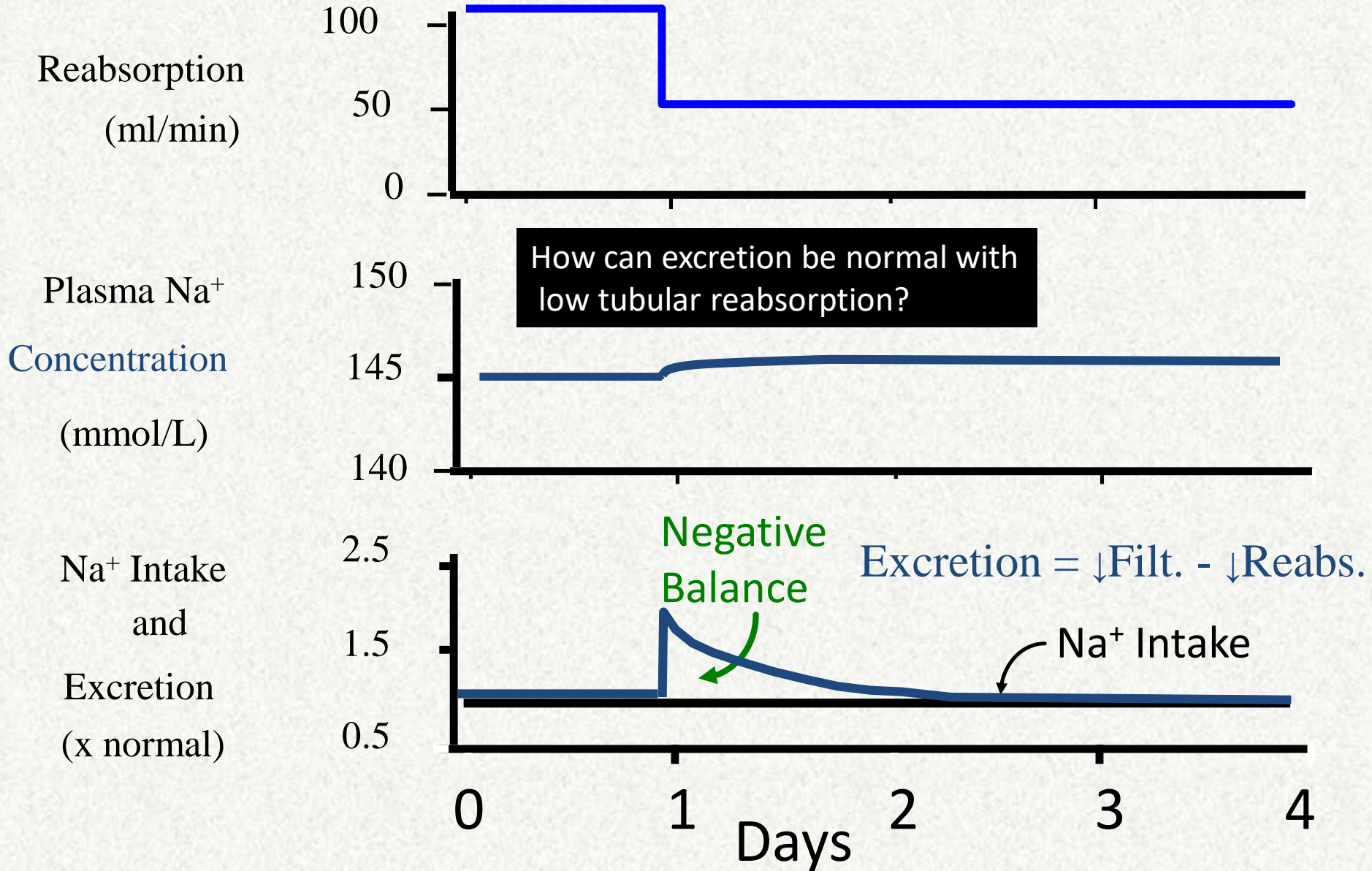
Glomerulo-tubular balance

Tubulo-glomerular feedback



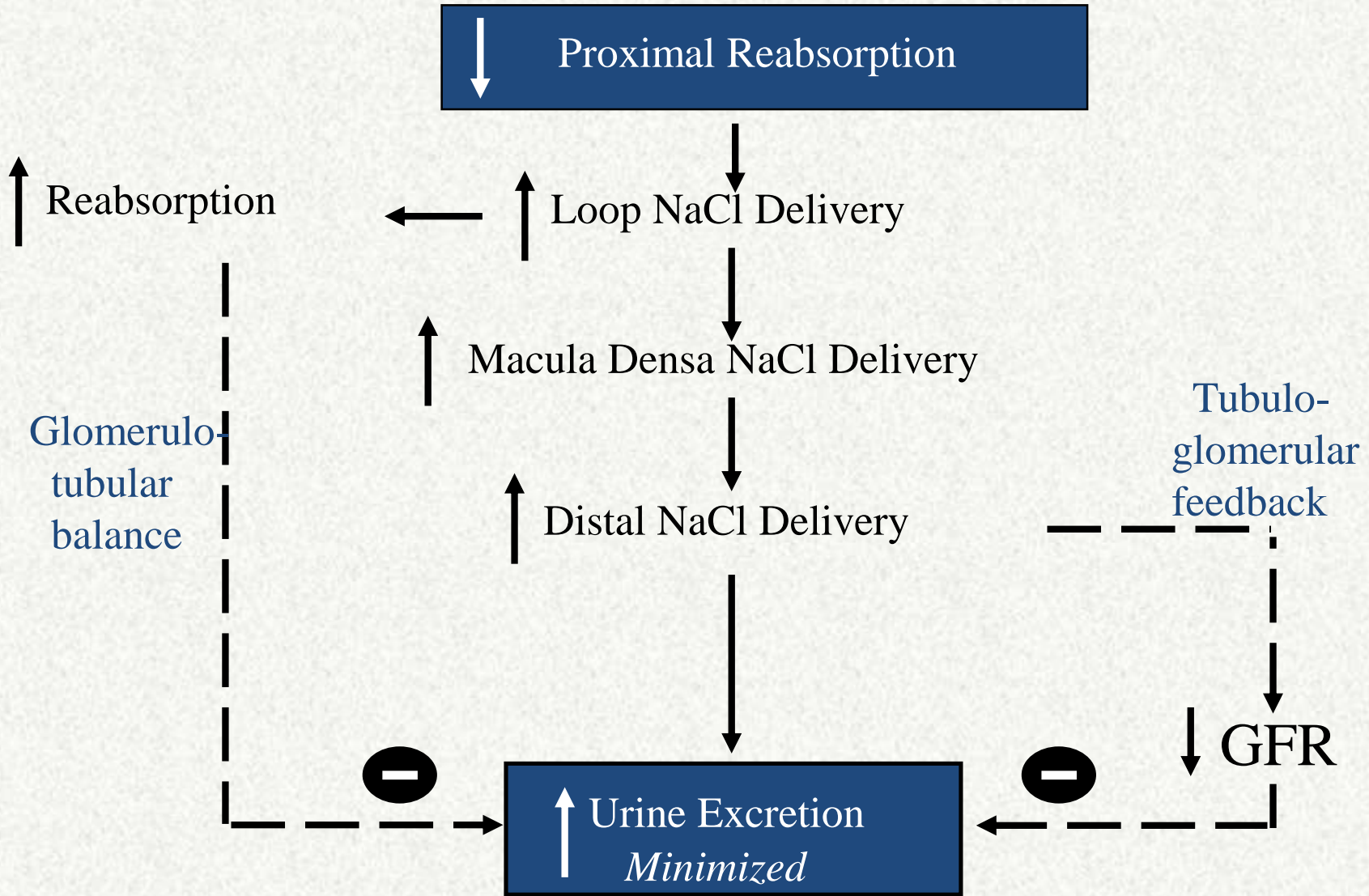


Effect of Decreased Reabsorption on Sodium Balance





Maintenance of Sodium Balance After Decreased Proximal Reabsorption





Hierarchy of Responses to Disturbances of Body Fluid Regulation

In steady-state, Intake = Output

1. Local renal responses

- changes in GFR
- changes in tubular reabsorption
- changes in tubular secretion

2. Systemic mechanisms (which can affect the whole body)

- changes in hormones
- changes in sympathetic activity
- changes in blood pressure
- changes in blood composition



Sodium excretion and extracellular fluid volume during diuretic administration.

Compensations that Permit Na^+ balance:

- \downarrow blood pressure
- \uparrow renin, angiotensin II
- \uparrow aldosterone

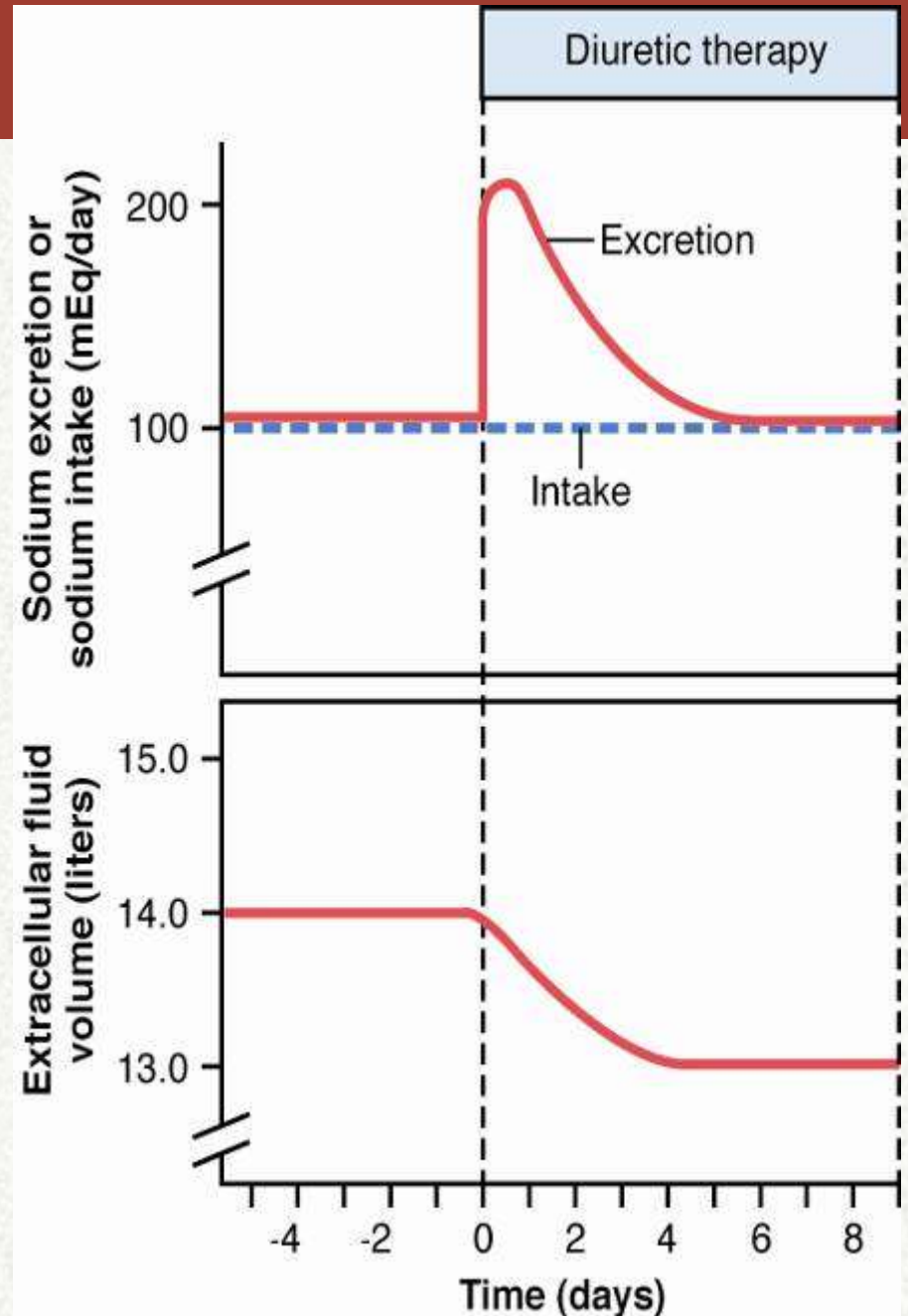
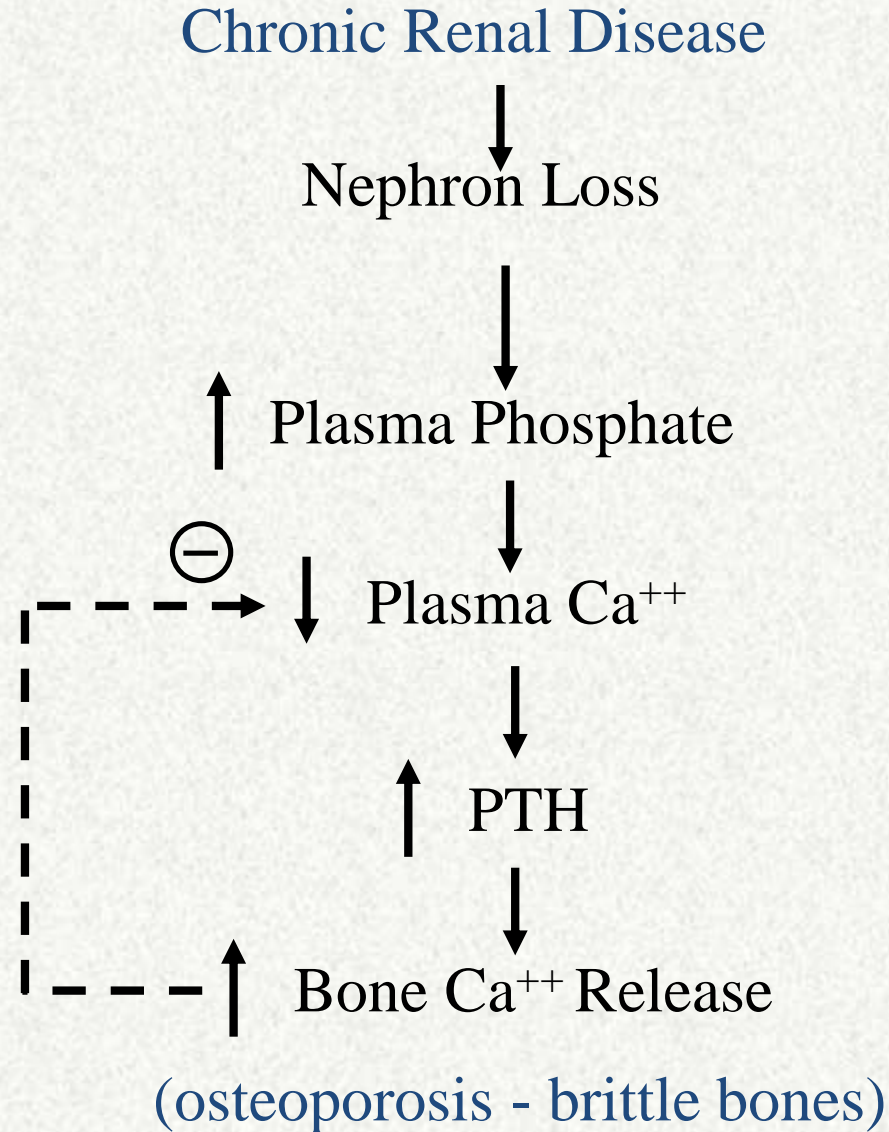


Figure 31-1



Hormonal Response to Chronic Renal Disease - PTH





Hierarchy of Responses to Disturbances of Body Fluid Regulation

In steady-state, Intake = Output

1. Local renal responses

- changes in GFR
- changes in tubular reabsorption
- changes in tubular secretion

2. Systemic mechanisms (which can affect the whole body)

- changes in hormones
- changes in sympathetic activity
- changes in blood pressure
- changes in blood composition



Renal-Body Fluid Feedback- Increased Fluid (Na^+) Intake

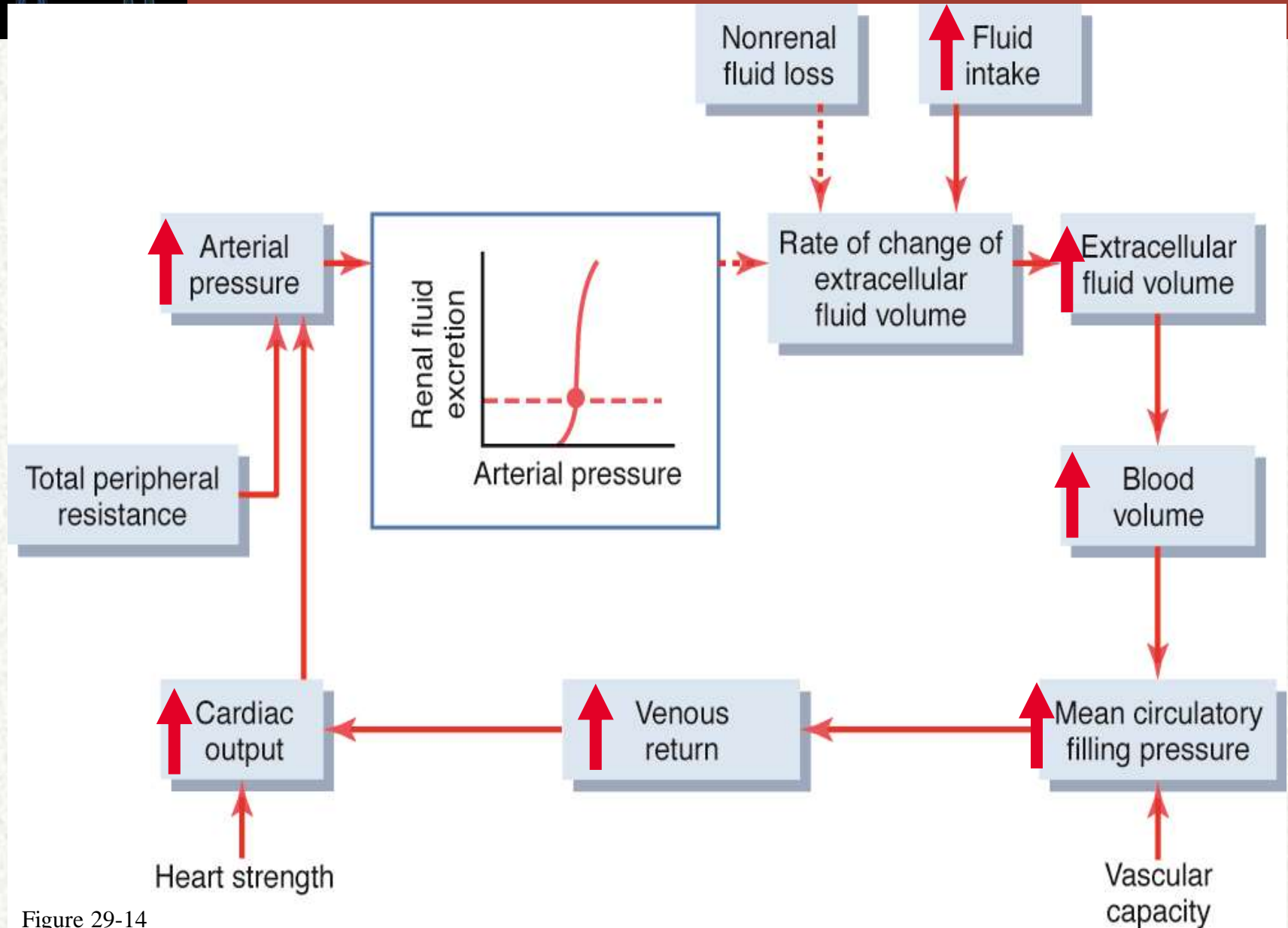


Figure 29-14



Integrated Responses to High Na⁺ Intake

Excretion Na⁺ = Filtration Na⁺ - Reabsorption Na⁺

1. Small increase in GFR
2. Decreased Na⁺ Reabsorption is caused by:
 - small increase in blood pressure
 - increased peritubular capillary pressure
 - decreased angiotensin II
 - decreased aldosterone
 - Increased natriuretic hormones (e.g. ANP)

Net effect = increased Na⁺ excretion