Tubulointerstitial Disease

Mark D. Baldwin D.O. FACOI
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Disclosures

• None, just working for The Man
Features of Tubulointerstitial Disease

1. Proteinuria - usually less than 1 Gm/da
2. Anemia - due to low level of Erythropoietin
3. Acidosis - RTA’s are common
4. Hypertension - common
5. Urinalysis - WBC’s and WBC casts seen
6. Electrolyte Abnormalities - Na and K

Many of the above features are seen at relatively mild elevations of Serum Creatinine
Features of Glomerular Disease

1. Proteinuria -> 3 Gm/da
2. Anemia - uncommon until late
3. Acidosis - uncommon until late
4. Hypertension - may occur at any time
5. Urinalysis - may see Oval Fat Bodies
6. Electrolytes - May see low Na

Unlike Tubulointerstitial Disease, many of these feature do not occur until late in the course of the underlying disease.
Types of Tubulointerstitial Disease

1. Acute Interstitial Nephritis
2. Chronic Interstitial Nephritis
3. Acute Tubular Necrosis
4. Renal Tubular Acidosis
5. Multiple Myeloma
Chronic Interstitial Nephritis (CIN)

• A chronic condition involving fibrosis of the interstitium and tubular destruction.
• Macroscopically normal kidneys
• The final common pathway of most chronic renal diseases
Causes of Chronic Interstitial Nephritis

- **Mechanical**: ureteral reflux, obstruction, stones, infection, neurogenic bladder, medullary cystic disease, Alport’s
- **Drugs**: NSAIDs, Lithium, PPI’s, cyclosporine, tacrolimus, indinavir, cisplatin
- **Heavy Metals**: Hg, Pb, Cd, arsenic, gold, uranium, too much Ozzy
- **Metabolic**: hyperuricemia, hypercalcemia, hypokalemia, hyperoxaluria, cytinosis
- **Radiation**
- **Immune mediated**: ANCA, SLA, Sjogren's, sarcoid
- **Vascular**: atherosclerotic renal disease
- **Heme/Onc**: myeloma, amyloid, lymphoma, sickle cell, PNH
- **Late glomerular disease**
- **Aristolochic acid**: Balkan nephropathy, Chinese herb nephropathy
Aristolochic acid

• **Acute exposure**: Chinese herb nephropathy-AKI, rapid decline in renal function

• **Chronic exposure**: Balkan endemic nephropathy-CIN/CKD slow decline over years from chronic exposure

Aristolochic acid Nephrotoxicity

Clinical Features of CIN

- Usually Asymptomatic
- STERILE PYURIA-The Hallmark of CIN
- Anemia
- Acidosis-Renal Tubular Acidosis
- Hypo or hyperkalemia
- Minimal Proteinuria
- Hypertension
Chronic Interstitial Nephritis
Chronic Interstitial Nephritis
Treatment of CIN

• Do NOT give antibiotics for pyuria unless there is bacteria present-this is a chronic inflammatory condition, NOT an infection
• BP control-The MOST important treatment
• ACE-I or ARBs-The drugs of choice
• Anemia control
• Acidosis control
• Phosphorus control
Renal Tubular Acidosis (RTA)

- **Distal Type I RTA** - associated with Chronic Urinary Tract Obstruction, Bicarb<15*NAG, hypokalemia, urine pH>5.5

- **Proximal Type II RTA** - associated with Fanconi’s Syndrome Bicarb 15-21*NAG, hypokalemia, urine pH>5.5

- **Distal Type IV RTA** - Most common RTA, Seen w/ DM and CKD, NAG, Hyperkalemia urine pH<5.5

- All RTA have +UAG

* Point of differentiation, NAG non anion gap acidosis
Urine Anion Gap (UAG) = (U_{Na^+} + U_{K^+}) - U_{Cl^-}

- NH_4^+ is the primary unmeasured cation which is not balanced by anions.
- UAG as indirect assay for renal NH4+ excretion
Urinary anion gap: \((\text{Na}^+ + \text{K}^+) - \text{Cl}^-\)

- In the presence of ammonium the chloride will be larger than the sum of Na and K.
- So a negative anion gap means ammonium in the urine.
- Ammonium in the urine means effective renal acid secretion
- Ammonium in the urine usually rules out RTA
Myeloma and the Kidney

- 10% of all hematological malignancies
- Plasma cell clone of Immunoglobulins usually IgG
- Renal, Cardiac and Liver are the most common organs involved
- Renal impairment - acute or chronic - is commonly seen
  ~50% of cases with severe involvement in 15-20% of cases
- Proteinuria - globulin or albuminurina is seen in >80% of cases
- Myeloma can involve the vascular, glomerular or
  tubular/interstitial segments of the kidney
- Frequent cause of mortality and morbidity
Pathophysiology

- Plasma cell clones leading to IgG light chains, heavy chains can be seen
- Can see clonal IgA, D, M or E variants of myeloma
- Light chain or fragments deposited in a tissues
- Kappa or lambda light chains
- Amyloid (AL) can be deposited
- Tubular obstruction
- Tubular dysfunction- Fanconi’s Syndrome & Proximal Type 2 RTA
- AKI of multiple etiologies
Renal Effects

- **Glomerular**: Amyloid light chain (AL) or heavy chain (AH) amyloidosis, Light Chain Deposition Disease (LCDD) or Heavy Chain Deposition Disease (HCDD) plasma cell infiltration

- **Tubular**: Cast nephropathy “Myeloma kidney”, tubular dysfunction, hypercalcemia, hyperuricemia, contrast induced AKI

- **Interstitial**: Plasma cell infiltration, pyelonephritis
Amyloid MM v LCDD

- **Amyloid myeloma** (AL):
  - lambda > kappa, + Congo red, + fibrils on EM

- **Light chain deposition disease** (LCDD):
  - kappa > lambda, - Congo red, - fibrils on EM
Cast Nephropathy

- Most common cause of renal failure in myeloma
- Globulin light chains are filtered at the glomerulus
- Can exceed 10-20 grams/day and are toxic to the tubular cells, negative dipstick d/t globulins not albumin
- Light chain are partially reabsorbed damaging to proximal tubular cells and delivered distally, combing with Tamm-Horsfell protein produced in the thick ascending limb occluding the tubule
- Obstructing casts lead to inflammation, fibrosis and tubular rupture
Presentation

- >50 years old
- Males > females
- African Americans > other groups
- Long history of back pain or “arthritic” pain
- Pathological fractures
- Fatigue
- Anemia
- Infection
- Renal failure
Multiple Myeloma Laboratory Findings

- Elevated BUN and Creatinine
- Hypercalcemia
- Hyperuricemia
- Normocytic Normochromic Anemia, Rouleaux formation
- Serum Protein Electrophoresis/ Urine Protein Electrophoresis positive for elevations in the Gamma fraction-M spike
- Low Anion Gap
- Urinalysis may show NO Protein, unless Sulfa salicylic Acid test is done, which will be positive.
Serology

- Serum protein electrophoresis/urine protein electrophoresis
- Free light chain assay
- Immunofixation: quantifies IgA, IgD, IgE, IgM, IgG
- Cytogenic analysis: karyotyping
- Flow cytometry
Congo Red “Apple green birefringence”
Renal Myeloma

Glomerular Amyloid

Cast Nephropathy
Amyloid Fibrils EM
Radiographs
Treatment of Myeloma Renal Disease

- Assure hydration status with alkalization of the urine but avoid fluid overload
- Allopurinol
- Bortezomib-dexamethasone-cyclophosphamide or Bortezomib-thalidomide-dexamethasone can decrease light chain production and may improve cast nephropathy
- Dialysis if needed (poor outcomes)
- ?role of plasmapheresis
Malignancy Related Hypercalcemic Renal Failure

- Prostate
- Renal Cell
- Breast
- Lymphoma/Leukemia
- Lung
- Myeloma
- Thyroid
- Other malignancies
Mechanism of Hypercalcemia

1. **Humoral Hypercalcemia of Malignancy** (HHM) 80%:
   - PTH related protein (PTHrP)
   - 1,25 dihydroxy Vit D (also seen w/ sarcoid and T.B.)
   - PTH-like substance (very rare)

2. **Osteolytic Metastasis**: 20%
   - Bone mets stimulate osteolysis via Osteoclast Activating Factor, RANKL, II, VEGF, TNF, TGF and PTHrP

3. **Prostaglandin mediated**
Presentation

- Decreased cognition, fatigue
- Anorexia, N, V, constipation
- Abdominal and bone pain
- Pancreatitis
- Short QT, ST changes pseudo MI pattern
- HTN
- AKI, Nephrogenic DI

Treatment of Hypercalcemia of Malignancy

- **Normal Saline**: Restores BP, decreases Ca reabsorption
- Calcitonin
- Bisphosphonates (Zolendronic acid most potent)
- Denosumab: Ab to RANKL, decreases OAF, best in cases refractory to Bisphosphonates
- **Loop diuretics**: enhance Ca excretion, don’t give thiazides will increase Ca reabsorption
Tubulointerstitial Diseases - Conclusions

- Often overlooked as a cause of Chronic Renal Disease
- Look for Drug causes or Sepsis as a cause of Acute Renal Failure (i.e. AIN or ATN)
- Tubulointerstitial Diseases frequently have electrolyte abnormalities, acid-base disorders, and anemia as a common feature.