Pathology Lecture - 2 Nephrotic Syndrome



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# The Nephrotic Syndrome

- a clinical complex resulting from glomerular disease & includes the following signs and symptoms:
- (1) massive proteinuria (3.5 gm /day in adults).
- (2) hypoalbuminemia ( $\leq 3 \text{ gm/dL}$ ).
- (3) generalized edema
- (4) hyperlipidemia and lipiduria.
- (5) little or no azotemia, hematuria, or hypertension.





## **Causes of Nephrotic Syndrome**

- 1- Primary Glomerular Diseases
- 2- Secondary (Systemic Diseases with Renal Manifestations)

Primary Diseases that Present Mostly with Nephrotic Syndrome

- 1- Minimal-change disease
- 2- Focal segmental glomerulosclerosis (FSGS).
- 3- Membranous nephropathy
- 4- membranoproliferative GN type 1 (usually a combination of nephrotic/ nephritic syndrome)

**Causes of Nephrotic Syndrome** 

### **1-primary glomerular diseases**

Cause	Prevalence (%) Children	Prevalence (%) Adults
Primary Glomerular Disease		
Membranous GN	5	30
Minimal-change disease	65	10
Focal segmental glomerulosclerosis	10	35
Membranoproliferative GN	10	10
IgA nephropathy	10	15

**Causes of Nephrotic Syndrome** 

#### **B-Systemic Diseases with Renal Manifestations:**

- Diabetes mellitus:
- Amyloidosis
- Systemic lupus erythematosus
- drugs (gold, penicillamine, "street heroin")
- Infections (malaria, syphilis, hepatitis B, HIV)
- Malignancy (carcinoma, melanoma)
- Miscellaneous (e.g. bee-sting allergy)

#### **1- Minimal-Change Disease (Lipoid Nephrosis)**

- benign disorder.
- The most frequent cause of the nephrotic syndrome in children (ages 1-7 years).
- <u>Pathogenesis:</u> still not clear.
- ? T-cell derived factor that causes podocyte damage and effacement of foot processes.



**Minimal change** disease. glomerulus appears normal, with a delicate basement membrane R diffuse effacement of foot processes of podocytes with no immune deposits.

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### **Morphology**

- <u>LM</u>
- the glomeruli appear normal.
- <u>IF</u>
- negative
- <u>EM</u>
- uniform and diffuse effacement of the foot processes of the podocytes .
- No immune deposits

#### MCD-EM

the capillary loop in the lower half contains two electron dense RBC's. Fenestrated endothelium is present and the BM is normal.

The overlying epithelial cell foot processes are fused (arrows).



# **MCD- Clinical Course**

- **nephrotic syndrome** in an otherwise healthy child.
- no hypertension.
- renal function preserved
- selective proteinuria (albumin)
- prognosis is good.
- Treatment: corticosteroids (90% of cases respond)
- < 5% develop chronic renal failure after 25 years
- In Adults with minimal change disease the response is slower and relapses are more common.

#### 2- Focal and Segmental Glomerulosclerosis (FSGS)

- sclerosis affecting some but not all glomeruli (focal involvement) and involving only segments of glomerulus.
- Usually nephrotic syndrome.
- It can occur :
- 1- as a primary disease( 20% to 30% of NS):
- e.g. inherited or congenital forms resulting from mutations affecting nephrin
- 2- Or: in association with underlying condition:
- e.g.; AIDS; heroin abuse; nephron loss; etc....

focal and segmental glomerulosclerosis (PAS stain).

a mass of scarred, obliterated capillary lumens with accumulations of matrix material (collagen)



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#### **MCD versus FSGS**

	MCD	FSGS
hematuria	-	+
hypertension	-	+
proteinuria	selective	nonselective
response to corticosteroid therapy	good	poor

- Pathogenesis
- unclear
- ? Genetics
- entrapment of plasma proteins and lipids in foci of injury where sclerosis develops.
- <u>Clinical Course</u>
- about 50% of individuals suffer renal failure after 10 years
- Poor responses to corticosteroid therapy.
- Adults do worse than children

### • <u>Morphology</u>

- LM:
- Sclerosis in some glomeruli not all of them; and in a segment not all of the affected glomerulus
- IF microscopy
- <u>Negative</u>
- **EM**
- effacement of podocyte foot processes

#### FSGS blue = collagen deposition (MT stain).



# **Collapsing glomerulopathy**

- a morphologic type of FSGS.
- poor prognosis.
- collapse of glomerular tuft and podocyte hyperplasia.
- It may be :
- 1-idiopathic .
- 2-associated with **HIV infection**.
- 3-drug-induced toxicities.

### **3-** Membranous nephropathy:

• Immune complex deposition in glomerulus

- <u>Types of Membranous glomerulonephritis :</u>
  1-Primary (85% of cases): antibodies against podocyte antigen phospholipase A2 receptor (PLA2R) antigen
- 2-Secondary to another condition or disease

**Secondary Membranous glomerulonephritis :** 

- (1) infections (HBV, syphilis, schistosomiasis, malaria).
- (2) malignant tumors (lung, colon and melanoma).
- (3) autoimmune diseases as SLE .
- (4) inorganic salts exposure (gold, mercury).
- (5) drugs (penicillamine, captopril,NSAID).

- Morphology
- LM
- diffuse thickening of the GBM .
- **IF**
- **deposits** of immunoglobulins and complement along the GBM (mainly IgG)
- **EM**
- subepithelial deposits "spike and dome" pattern.



Membranous nephropathy. subepithelial deposits and the presence of "spikes" of basement membrane material between the immune deposits .

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# A silver stain (black). Characteristic "spikes" (green arrows) seen with membranous glomerulonephritis as projections around the capillary loops.



Membranous GN IF: granular deposits of IgG and complements along the capillary walls



#### EM: "spike and dome" pattern is characteristic Dome= immune complex spike= basement membrane



- <u>Clinical Course</u>
- nephrotic syndrome
- poor response to corticosteroid therapy.
- 60% of cases → proteinuria persists
- ~40%→ progressive disease and renal failure within 2 to 20 yr.
- 30% →partial / complete remission of proteinuria.