Systematic pathology



Kidney and urinary tract pathology

Kidney diseases





Congenital diseases

⇒ Adult polycystic kidney disease

• common congenital disease, ↓of renal function in the 3.- 4. dec., autosomal dominant - gen usually on the short arm of chromosome 16

• gross: symmetrical kidney enlargement – lenght to 30 cm, multiple cysts 0,5-50mm





⇒ Solitary kidney cysts

accidental finding . Important diff. dg x cystic renal carcinoma

Polycystic kidney





Vascular kidney disorders



- Renal artery stenosis
 - renovascular hypertension (Goldblatt's)
 - pressure \(\) in afferent arterioles
 - \Rightarrow \downarrow of filtration pressure in the glomerulus
 - juxtaglomerular apparatus hyperplasia + renin overproduction
 - blood pressure \(\frac{1}{2}\) by longer duration vascular atrophy.

Vascular kidney disorders

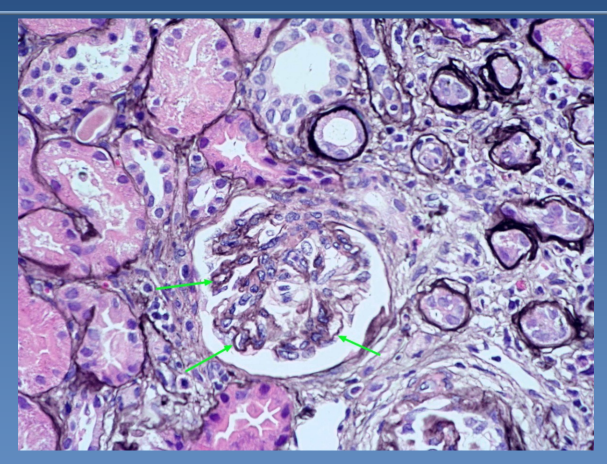


Benign nephrosclerosis

- > by benign (compensated) hypertension
 - gross: symmetrical decrease in size, fine granulated surface
 - micro: hyalinne insudates in arteriolar walls, median hypertrophy + intimal sclerosis, ischemic changes +/glomerular loss, vascular atrophy of the tubules, adjacent interstitial fibrosis.



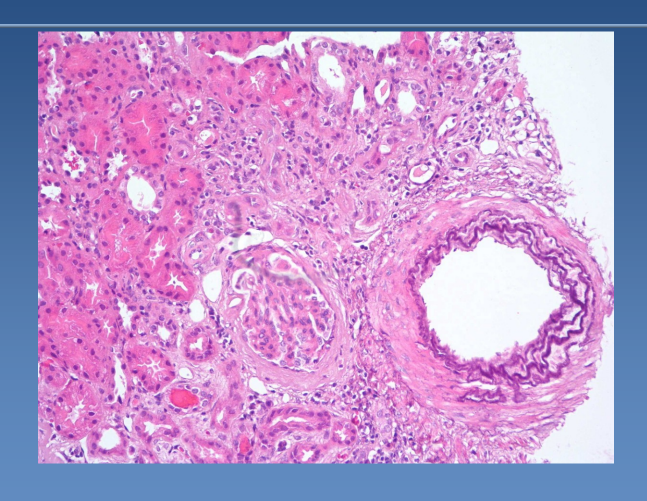
Benign nephrosclerosis



Ischemic glomerular chamges, "wrinkling" of the GBM

Benign nephrosclerosis





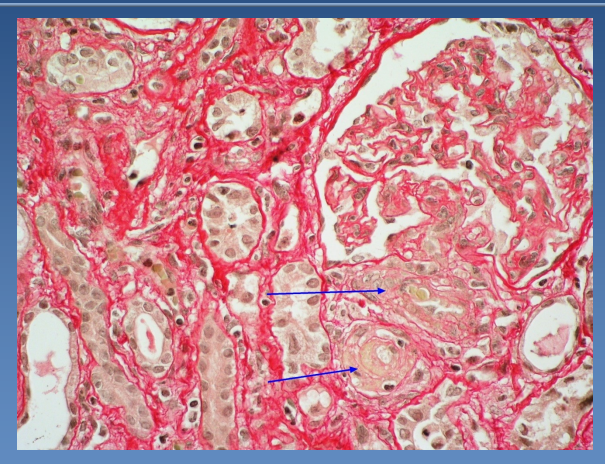




- Malignant nephrosclerosis
 - due to accelerated arterial hypertension (diastole >130mmHg), endothelial damage
 - •gross: renal oedema, infarctions possible
 - micro: oedema, intimal mucoid seepage in arteries,
 fibrinoid necrosis of the arteriolar wall, possible trombi

Malignant nephrosclerosis

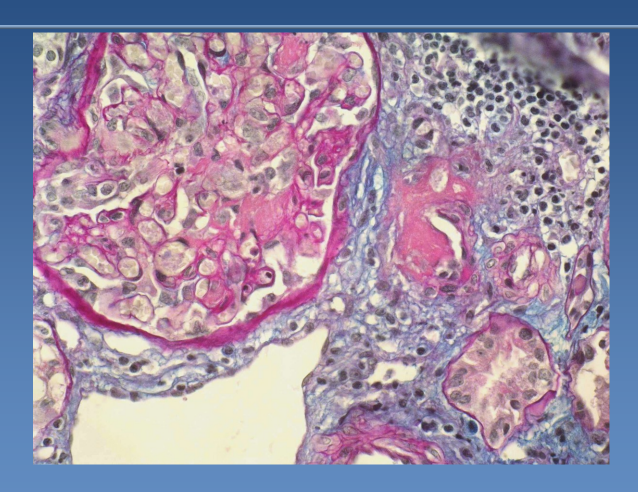




Singificant arteriolar luminal narrowing, endothelial oedema



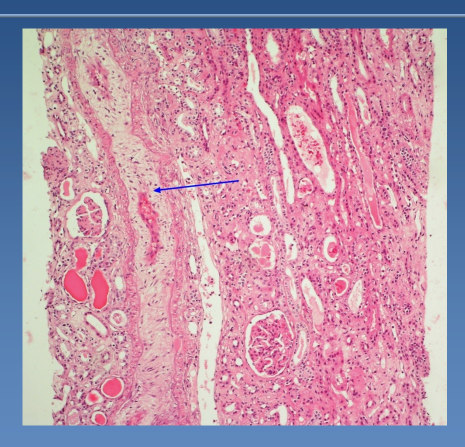


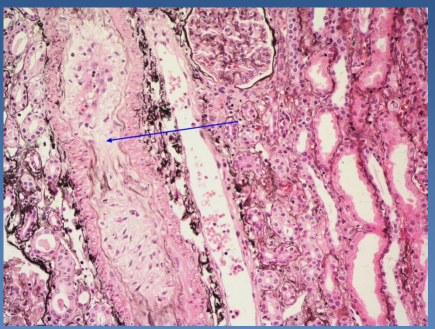


Fibrinoid necrosis of the hilar arteriole

Malignant nephrosclerosis







Oedema, mucoid intimal seepage, luminal narrowing in a muscular artery

Vascular kidney disorders



- renal infarction
 - ischemic coagulative necrosis due to blockage of peripherale branches of the renal artery
 - •gross: yellowish conical necrosis
 - micro: necrosis with haemorragic rim

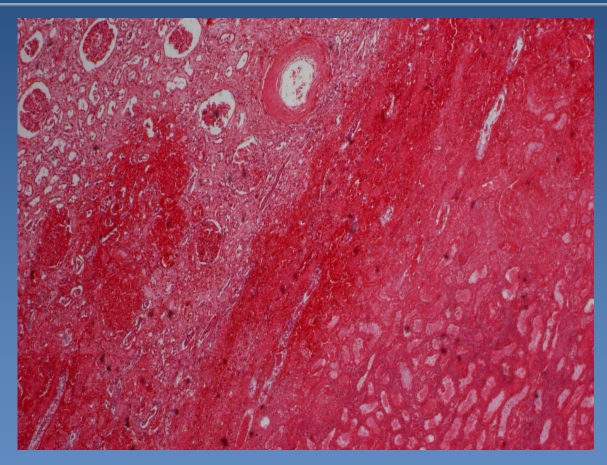
Renal infarction







Renal infarction



coagulativeí necrosis

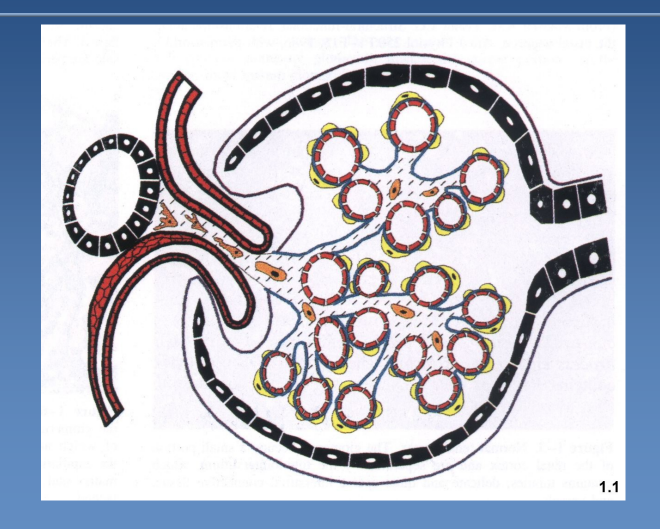
Glomerular diseases



- Glomeral damage caused by various factors
 - vascular changes
 - metabolic diseases
 - familiar diseases
 - immune-mediated disorders







Mechanism of the glomerular damage

Immune-mediated damage

- circulating immune complexes
- in situ immune complexes
- anti-GBM antibodies
- antineutrophilic antibodies

Mechanism of the glomerular damage

Non-immunological damage

- haemodynamic factors
- hypertension
- ischemia

Glomerular reaction to the damage



proliferation:

⇒hyperplasia of mesangial, endothelial, epithelial cells – hypercellularity. Epithelial cells (podocytes) may be a part of crescents filling the Bowman's capsule.

- ***** exudation:
- leukocytes + fibrin
- * thickening of the glomerular capillary wall
- usually due to deposition of immune complexes and/or GBM reaction





***** sclerosis:

eosinophilic material consisting of the mixture of collapsed membranes, mesangial matrix and plasmatic proteins. PAS + silver impregnation highly positive

hyalinosis:

foci of refractive amorphous material comprising insudated plasmatic proteins and lipoproteins (PAS intensive positivity, silver impregnation negative)

Clinical presentation of the glomerular disorders

- According to the number of affected glomeruli
 - diffuse changes (> 50% of gl.)
 - focal changes

- According to the extent of glomerular lesion
 - global changes (the whole gl.)
 - segmental changes

Clinical presentation of the glomerular disorders

- nephritic syndrome:
- acute gl. damage, hematuria, proteinuria, oligouria, oedema, hypertension

- nephrotic syndrome:
- severe proteinuria with protein loss > 3,5g/24h, hypoalbuminemia, decrease of production of concentrated urine, oligouria \rightarrow anuria, \uparrow azotemia

Clinical presentation of the glomerular disorders

- acute renal failure:
- ⇒ sudden decrease of production of concentrated urine, oligouria → anuria, ↑ azotemia
- chronic renal failure:
- gradual loss of renal functions

Glomerular diseases classification



Mostly according to the clinical signs

Glomerulopathy with proteinuria or nephrotic syndrome

Glomerulopathy with isolated or predominant hematuria

Glomerular diseases classification



Glomerulopathy with acute nephritic syndrome

Glomerular/kidney involvment by SLE

Chronic glomerulonephritis



Glomerular diseases classification

- primary x secondary GN
 - ⇒ primary GN disorder limited to the kidney, without systemic disease
 - secondary GN part of other disease (SLE, hepatitis C, neoplasia, ...)

Glomerulopathies manifestated by proteinuria/nephrotic sy

Proteinuria with nephrotic syndrome

Minimal change disease

Focal segmental glomerulosclerosis

Membranous glomerulopathy

Amyloidosis

Diabetic nephropathy

Glomerulopathies with proteinuria/nephrotic sy

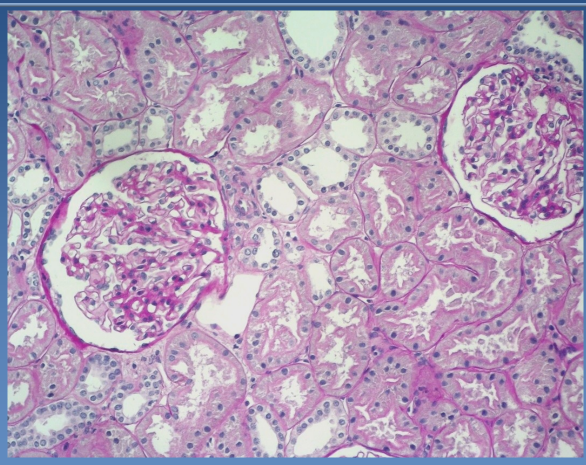


- Minimal glomerular change disease
- mostly in children's age
- ⇒ heavy selective proteinuria (albuminuria)
- nefrotic syndrome responsive to steroid therapy
- normal renal functions

- LM: normal glomerular morphology
- IMF: negative, without immunodeposits
- EM: diffuse fusion of podocytes' foot processes

Minimal glomerular change disease

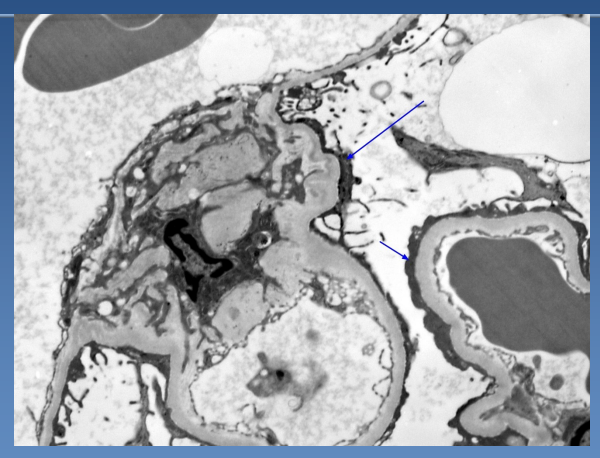




Normal glomerular morphology

Minimal glomerular change disease (EM)





diffuse fusion of podocytes' foot processes

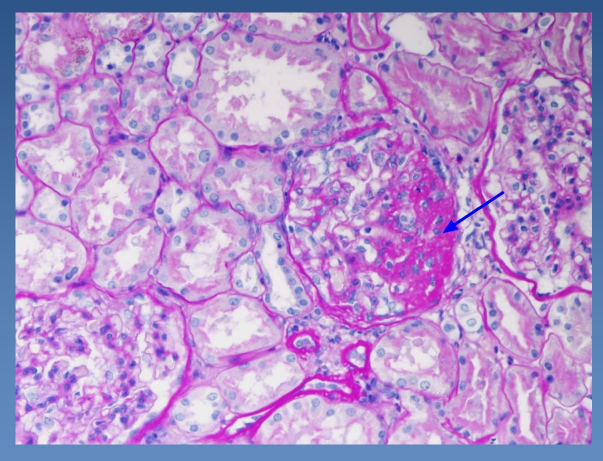
Glomerulopathies with proteinuria/nephrotic sy



- Focal segmental glomerulosclerosis (FSGS)
 - ⇒ children, adults (↑ incidence)
 - non-selective proteinuria up to nephrotic type
 - nephrotic syndrome, steroid-resistant
 - gradual progression to the renal failure
 - LM: Focal segmental sclerotic and hyalinne gl. changes due to capillary loops collapse and mesangial expansion
 - IMF: negative, without immune deposits
 - EM: fusion of podocytes' foot processes and podocytes' detachment from the GBM

FSGS





Segmental sclerosis of the capillary tuft

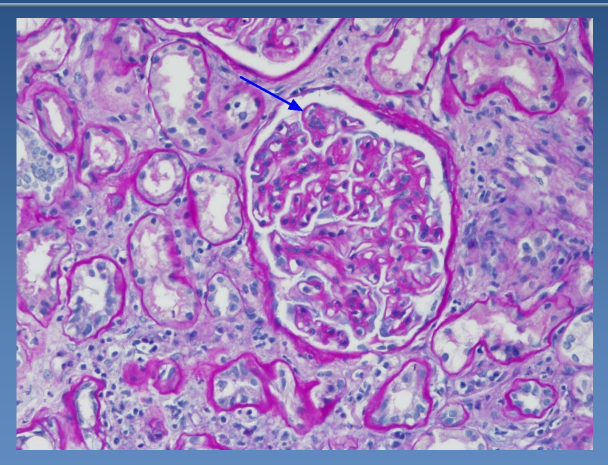
Glomerulopathies with proteinuria/nephrotic sy



- Membranous glomerulopathy
 - immune complex-mediated glomerulopathy, mostly in adults.
 - proteinuria of nephrotic type, hematuria.
 - LM: diffuse and global gl. involvment, normocellular. Deposition of immune complexeson the outer aspect of the glomerular basal membrane (GBM), thickened in futher stages.
 - IMF: granular deposits along GBM (IgG, C3)
 - EM: subepithelial electron-dense immune deposits



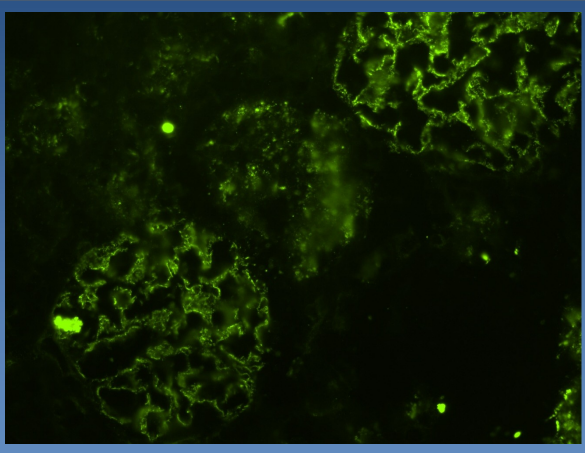
Membranous glomerulopathy



Diffuse GBM thickening
Glomerulus without inflammation or proliferation

Membranous glomerulopathy (IMF)

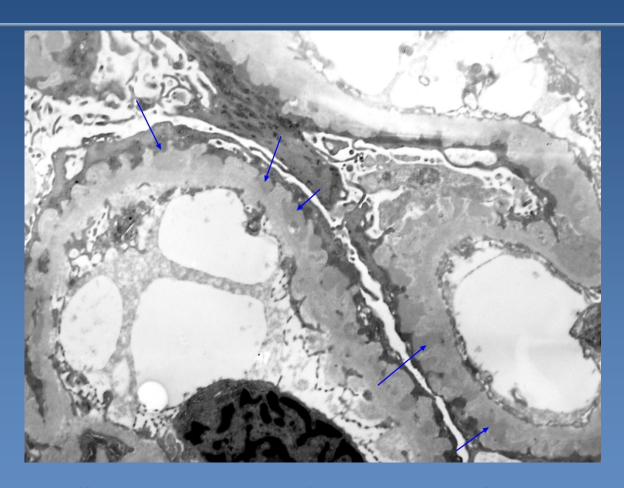




Granular deposits along the GBM in IgG

Membranous glomerulopathy (EM)





Diffuse subepithelial (outer aspect of the GBM) immune deposits

Glomerulopathies with proteinuria/nephrotic sy



- extracellular deposition of pathological fibrilary protein with typical staining features
- > systemic amyloidoses most clinically important
- 4 main groups:
 - AA amyloidosis (SAA protein precursor) in chronic diseases (RA, IBD, ...)



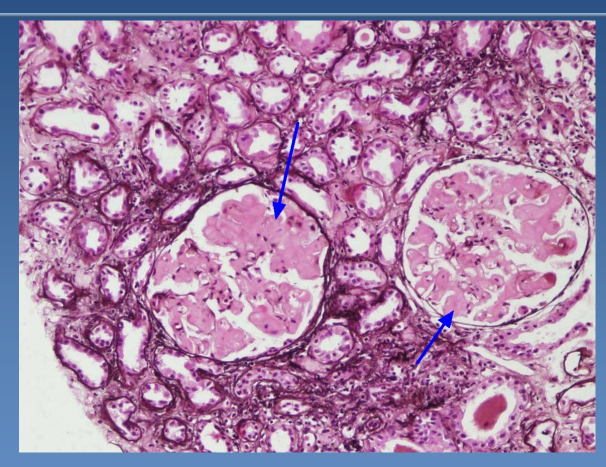
- AL amyloidosis (precursor plasma cell product) in monoclonal plasma cell disorders
- hereditary amyloidosis: genetically determinated protein defect (transthyretin)
- amyloidosis associated with haemodialysis





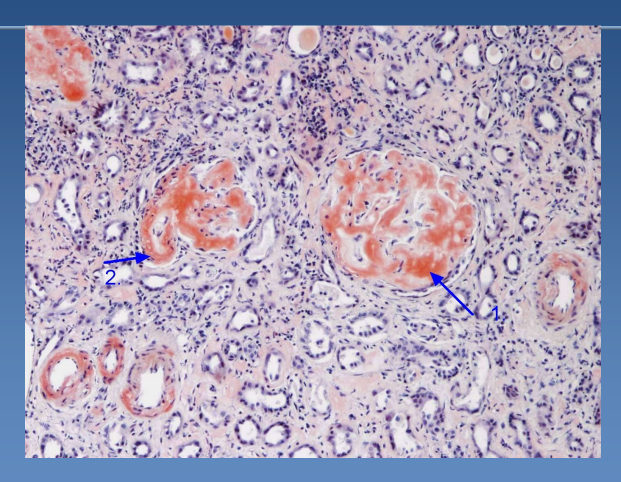
- proteinuria with nephrotic syndrome
 - LM: structure-less eosinophilic masses in the glomeruli, tubules, intersticium and vessels
 - Positive Congo red staining, green dichroism in polarisation
 - IMF: positivity of AA amyloid, light chains
 - EM: non-branching, randomly orientated fibrils, size of 6-13nm





Amyloid deposition in the glomerulus





Congo red-positive amyloid deposition in the glomerulus

Glomerulopathies with proteinuria/nephrotic sy



- Diabetic glomerulopathy
 - renal involvment by diabetic microangiopathy
 - proteinuria of nephrotic type
 - LM: thickening of GBM, mesangial expansion by PAS positive mesangial matrix, mildly increased cellularity, glomerular enlargement – diffuse diabetic glomerulosclerosis





 later homogennous eosinophilic nodular formations, mesangial cells pushed to the periphery – nodular diabetic glomerulosclerosis.

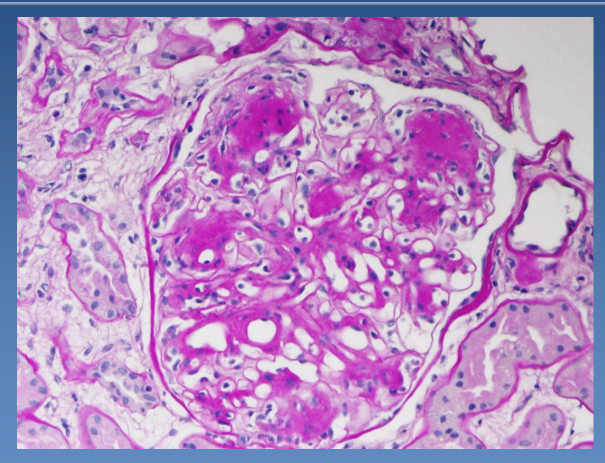
Hyalinne insudations in arterioles

•IMF: without immune deposits

•EM: thickening of GBM



Diabetic glomerulopathy



Mesangial nodules



Glomerulopathies with isolated or prevalent haematuria

IgA nephropathy (Berger's disease)

Henoch-Schönlein purpura

Alport syndrome / thin basement membranes sy



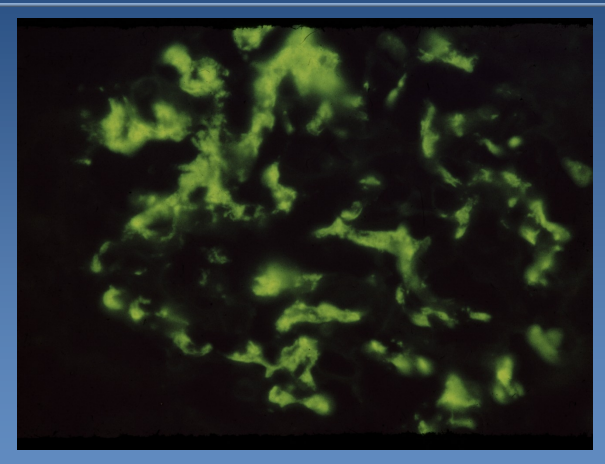
- IgA nephropathy (Berger's disease)
 - ⇒ immune complex-mediated disorder with raised levels of circulating IgA
 - ⇒ IgA mesangial deposits by chronic GIT, respiratory tract mucosal inflammations, liver cirrhosis
 - episodic macroscopic haematuria in coincidence with respiratory infection



- LM: mesangial proliferation
- IMF: mesangial IgA granules
- EM: Mesangial and paramesangial ID
- Henoch-Schönlein purpura
 - extensor skin vasculitis with purpuric rash, GIT manifestations, arthralgia
 - renal involvment IgA nephropathy

IgA nephropathy IMF





Mesangial IgA immune deposits



- Alport syndrome/ thin basement membrane lesion
 - ⇒ mutation in genes for collagen IV, part of basement membranes, (mostly gene COL4A5 encodedon the X. chromosome).
 - gradual progression of renal failure
 - in the fully evolved Alport sy − bilateral hearing disorders, ocular abnormities

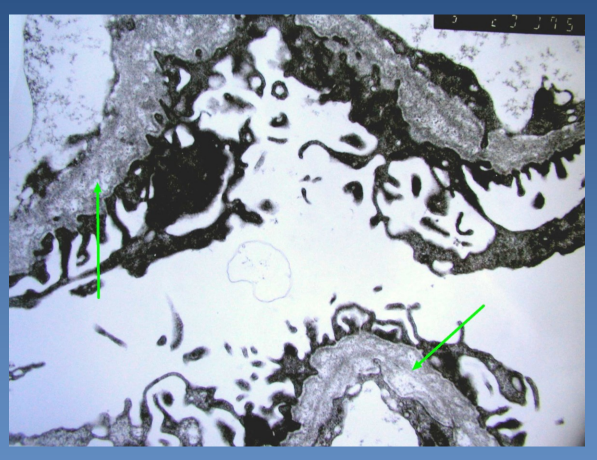
Alport syndrome/ thin basement membrane lesion

* thin basement membrane lesion

- without progression into renal failure, mild clinical signs (benign familiar haematuria)
- ⇒ typical morphology possible in female carriers of X-linked
 Alport syndrome

Alport syndrome/ thin basement membrane lesion ELMI





Characteristic picture of lamellar glomerular basement membrane in hereditary nephropathy.





Glomerulopathies with acute nephritic syndrome

Acute diffuse proliferative GN

Membrano-proliferative GN

Rapidly progressive glomerulonephritis (RPGN)

Glomerulopathies with acute nephritic syndrome



usually proliferative glomerulonephritis with increased mesangial and endocapillary cellularity, commonly with crescent formation.

- Acute diffuse endocapillary proliferative GN
 - syn. acute post-infective, acute proliferative, exudative GN
 - immune complex-mediated disorder

Acute diffuse endocapillary proliferative GN



- ⇒ usually post-infective glomerulonephritis (beta-hemolytic streptococcus, staph., G-bacteria, viruses, parasites)
- ⇒ systemic disorders (SLE, infective endocarditis, nectorising arteritis) may be accompanied by this GN
- most commonly children, 1-4 wks. after streptococcal infection



- ⇒ haematuria, proteinuria, hypertension, oedemas, renal failure
- possible asymptomatic course
- ⇒ raised ASLO titre and drop of C3, C4 complement in serum

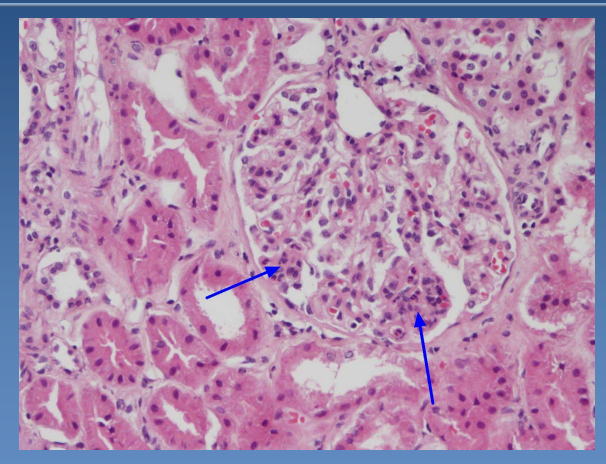


- benign course in children
- protracted course in adults, with hypertension, variable grade of renal failure
 - LM: ↑ endocapillary and mesangial cellularity, capillary lumen compression



- •IMF: diffuse segmental IgG and C3 granules in capillary loops, in mesangium
- •EM: humphs electron-dense subepithelial immune deposits

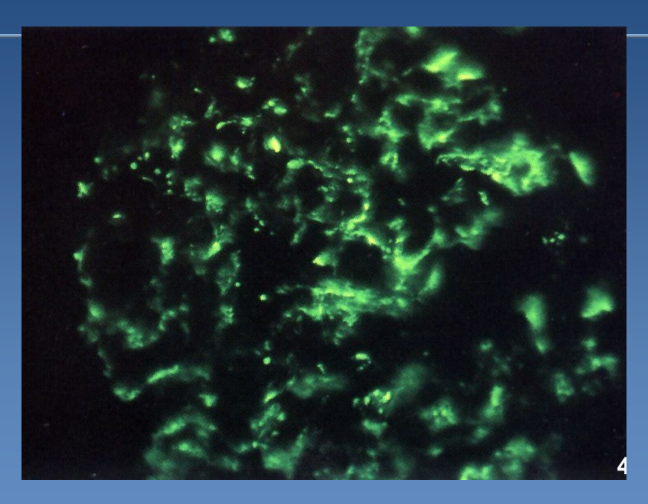




hypercellularity, neutrophils

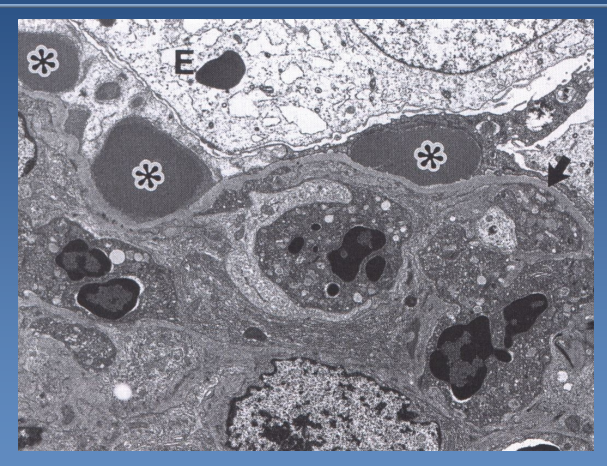
Acute post-infective GN (IMF)





Granular IgG deposits on GBM and in mesangium





Granular subepithelial deposits

Glomerulopathies with acute nephritic syndrome

- e
- → Membrano-proliferative GN (mesangio-capillary)
- Type I.-III. according morphology,
- commonly C3 glomerulopathy (problem in complement C3 activation control), activation of mesangial + endothelial cells
 - Type I. immune complex-mediated, cryoglobulinemia (esp. hepatitis
 C), other causes more common in children, teens
 - \(\) serum complement, nephritic syndrome, nephrotic sy possible.
 - LM: diffuse glomerulopathy, endocapillary and mesangial hypercellularity, accentuation of capillary tuft lobular architecture,
 GBM duplication ("tram track") in PAS, silver impregnation.

Membrano-proliferative GN



- •EM: subendothelial immune deposits + mesangial interposition (inclusion of mesangium + new layer of BM inbetween the immune deposits and original BM duplication, "splitting"), subendothelial + mesangial ID.
- ⇒ Type II. dense deposit disease
- \Rightarrow >60% of patients with antibody C3nephritic factor (NeFa) binding to C3 convertase \rightarrow stabilisation (no enzymatic degradation), \rightarrow permanent C3 activation of alternative pathway of complement cascade

Membrano-proliferative GN



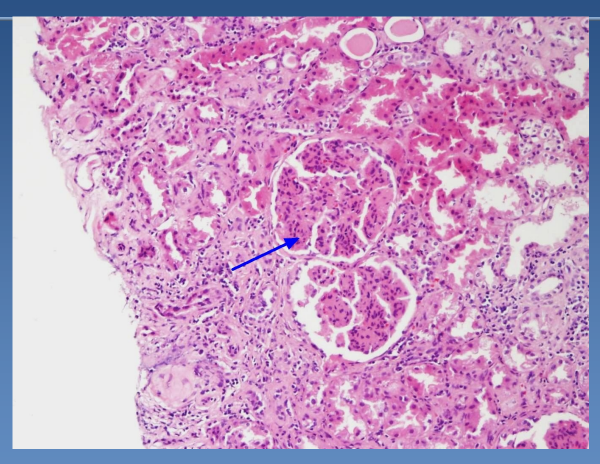
•EM: dense-deposit disease (DDD). Ribbon-like immune deposits in the GBM and mesangium,

⇒ Typ III. rare

- LM: same findings as in the type I.
- EM: + subepithelial ID.

Membrano-proliferative GN

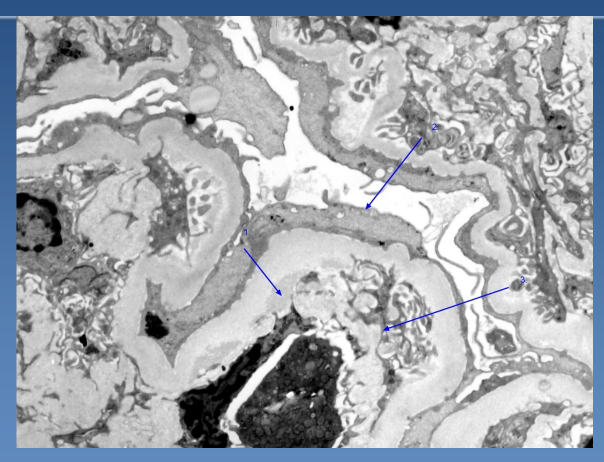




Lobulisation of the capillary tuft, hypercellularity in mesangium + endocapillary

Membrano-proliferative GN (EM)





- 1. Subendothelial immune deposits 2. podocyte foot processes fusion
- 3. mesangial interposition

Glomerulopathies with acute nephritic syndrome



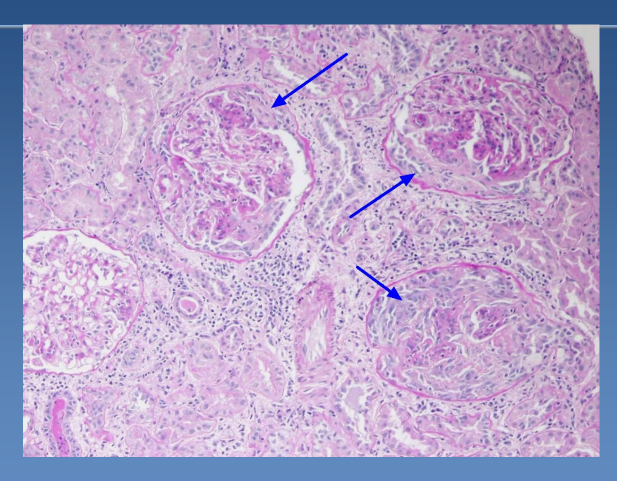
- Rapidly progressive GN (RPGN), crescentic
 - Hematuria, proteinuria
 - Rapid loss of renal functions
 - Extensive crescentic formation



- Variable group of diseases:
- ⇒ pauci-immune GN (part of systemic vasculitis, sm.
 ANCA+)
- ⇒ Anti-GBM disease
- ⇒ immune-complex mediated GN

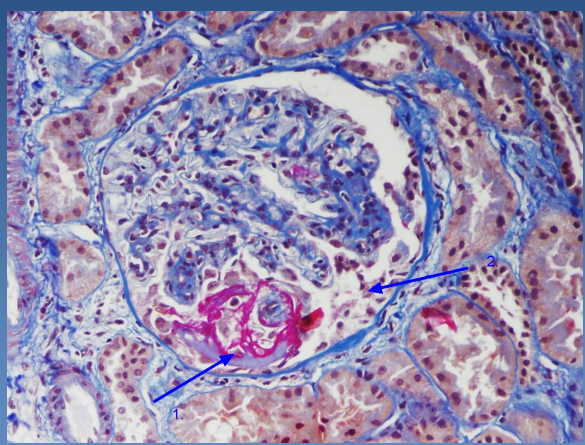
 complication of other GN (IgA, post-infectious GN, SLE)





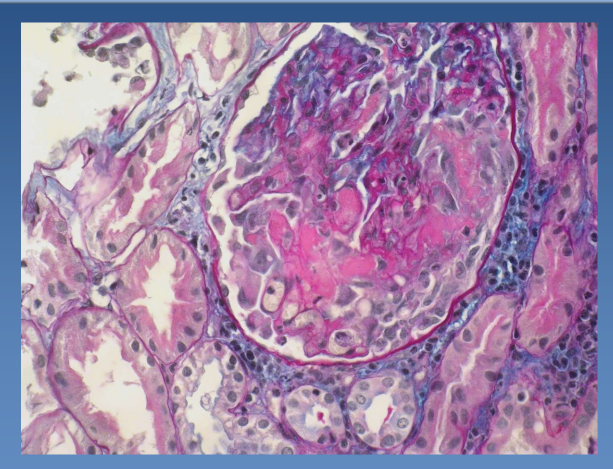
Cellular crescents within the Bowman capsule





- 1. Fibrin in the crescent
- 2. Cellular crescent (incipient)





Fibrinoid necrosis of capillaries

Systemic vasculitis

anti-GBM vasculitis immune-complex mediated vasculitis ANCA-associated vasculitis

Hypertensive kidney disorders

Thrombotic microangiopathy

Others

renal infarction renal artery stenosis



- Systemic vasculitis
- Anti-GBM glomerulonephritis
 - antibodies against Goodpasture antigen (part of noncollagenous portion of the GBM)



- \Rightarrow binding of anti-GBM antibody \rightarrow complement + proteases activation k aktivaci \rightarrow GBM destruction
- ⇒LM: RPGN appearance
 - •IMF: diffuse linear IgG deposits positivity of GBM

Immune complex-mediated vasculitis

- ⇒ Henoch-Schönlein purpura
 - IgA nephropathy morphology

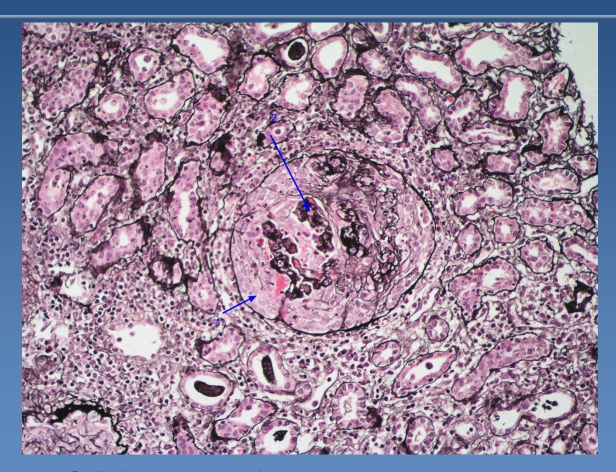


ANCA-associated vasculitis (antineutrophil cytoplasmic antibodies)

- ⇒ Granulomatosis with polyangiitis (Wegener granulomatosis)
- - RPGN morphology



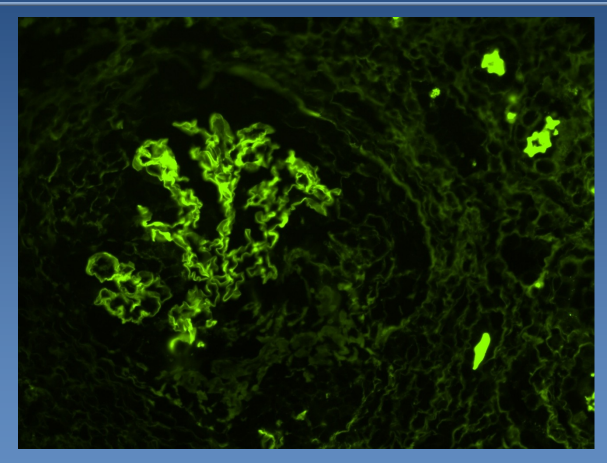
Anti - GBM



- 1. Cellular compressive crescent
- 2. Collapsing capillary tuft



Anti-GBM (IMF)



Linear peripheral IgG positivity (on the GBM)



Thrombotic microangiopathy

- → Haemolytic uremic syndrome, Thrombotic thrombocytopenic purpura – formation of platelet thrombi in small vessels of systemic circulation, platelets consumption, endothelial damage and haemolysis
- ➡ Intimal and endothelial oedema, fibrinoid necrosis of the arteriolar wall, fibrin thrombi in capillaries

>types:

- epidemic (E.coli shiga-like toxin)
- other drugs, irradiation, infection
- TTP hereditary/acquired excessive activation of platelets

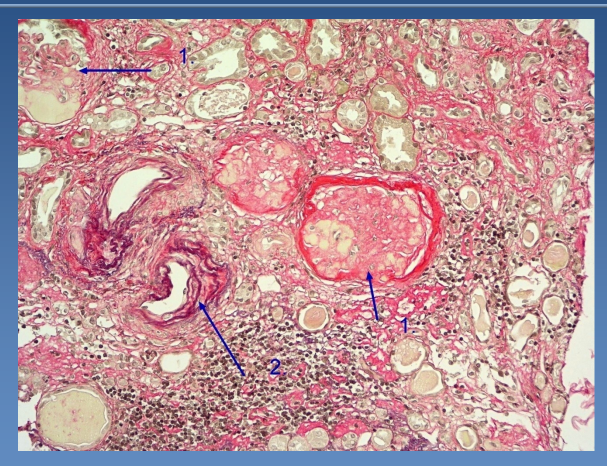
Chronic glomerulonephritis

gl. disease in the end-stage (significant renal lesion)

- gross: kidney contracted, granulated
- micro: high percentage of globally obliterated glomeruli, interstitial fibrosis, tubular atrophy, vascular changes.

Chronic glomerulonephritis





- 1. Obliterated glomeruli
- 2. Vascular changes

- both parts (tubules + interstitium) affected
- Two main categories:
 - ⇒ Ischemic and toxic lesion (acute tubular necrosis ATN)

Inflammatory (tubulointerstitial nephritis TIN)

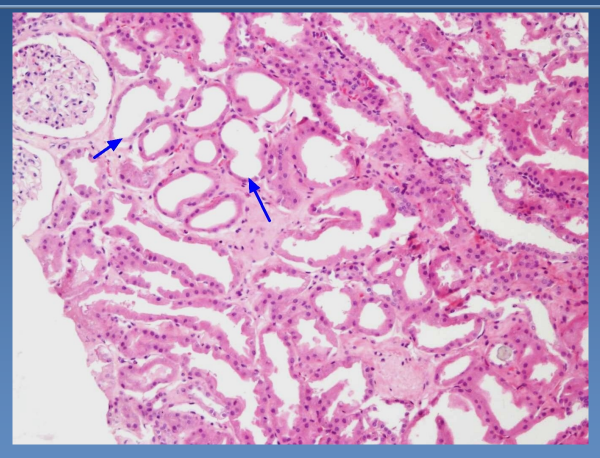


*Acute tubular necrosis

- etiology: ischemic, toxic
- ⇒ acute renal failure with oligouria/anuria, hemodialysis necessary
 - •gross: kidney edema, markedly pale cortex
 - micro: variable grade of tubular cells injury, from loss of brush border to necrosis.
- Ischemic segmental lesions along the whole tubular lenght
- toxic proximal tubules



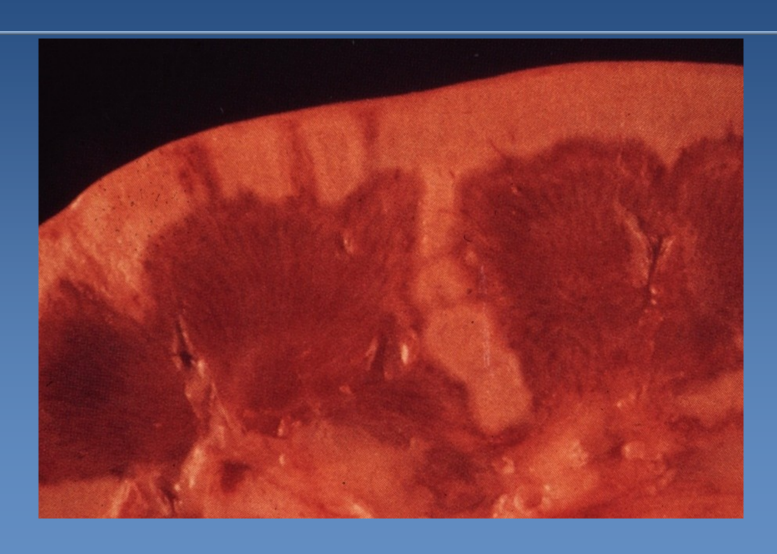
Acute tubular necrosis



Tubular dilatation, simple flat epithelium

Acute tubular necrosis







- * Acute tubulo-interstitial nephritis
 - Etiology: infectious bacterial (acute pyelonephritis)
 - toxic drug-induced (post ATB)
 - metabolic (diseases with crystal formation)
 - viral (hantaviruses)
 - micro: interstitial inflammatory infiltrate, variable grade of tubular epithelium injury

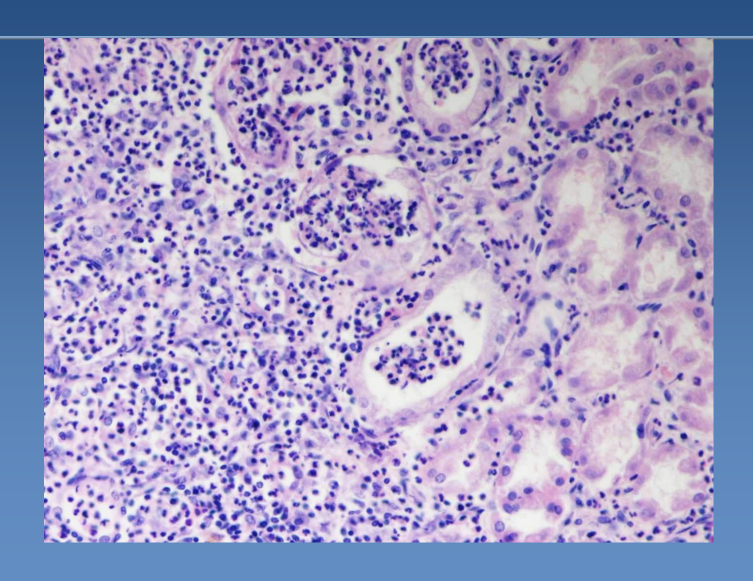


- Acute pyelonephritis
 - ⇒ acute pelvis + kidney inflammation mostly ascennding bacterial infection i.e. E. coli
 - descending in sepsis
 - febrile ilness, lumbal pain, dysuria + urging, pyuria with numerous neutrophils

- •gross: swollen kidney, yellow subcapsular abscesses.
- edematous, hyperemic pelvis, sm. with pus, progression of purulent inflammation to the adjacent tissues paranephritic abscess
- •micro: interstitial + tubular neutrophils

Acute pyelonephritis







- Chronic pyelonephritis
 - one of the most common causes of renal failure

possible insidious start, manifestation due to hypertension, commonly after multiple attacks of acute pyelonephritis.

- gross: irregular shrunken kidney, flat scars, commonly + nephrolithiasis, progressive atrophy end-stage kidney
- micro: interstitial fibrosis, tubular atrophy, dilatation
 + casts (follicular colloid-like), glomerular hyalinisation

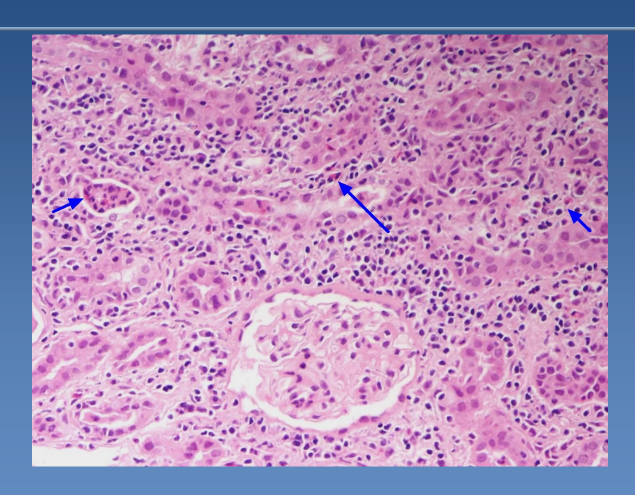


Drug-induced TIN

- Antibiotics, NSAIDs
 - micro: interstitial oedema, mixed interstitial inflammatory infiltrate with eosinophils



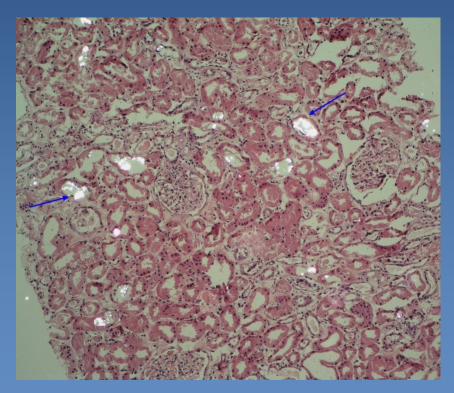


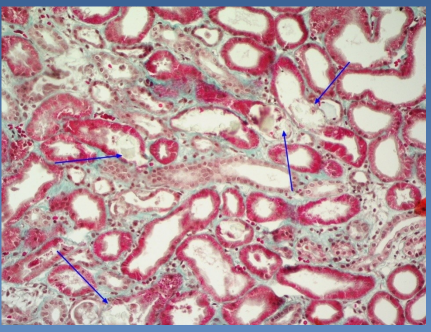


Eosinophils in inflammatory infiltrate

Oxalate nephropaty







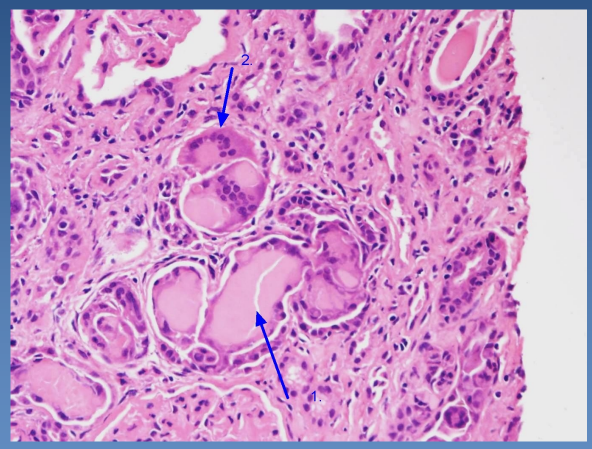
Oxalate crystals/deposits in tubules



- Myeloma nephropathy
 - renal damage due to myeloma
 - excretion of light chains (BJ protein) into primary urine, toxic to epithelia
 - → + casts formation → nephrohydrosis, blockage of urine outflow within renal parenchyme.
 - tubular epithelial damage, multinucleated macrophages

Myeloma nephropathy





- 1. Protein casts
- 2. Giant multinucleated macrophages

Renal tumors



- Benign x malignant
- ***** Benign
 - ⇒ angiomyolipoma
 - Mesenchymal (perivascular epithelioid cell PEComa)
 more common in patients with tuberous sclerosis
 - ⇒ cortical adenoma
 - •micro: papillary structure
 - •gross: ochre colour, size< 5mm</p>
 - accidental finding



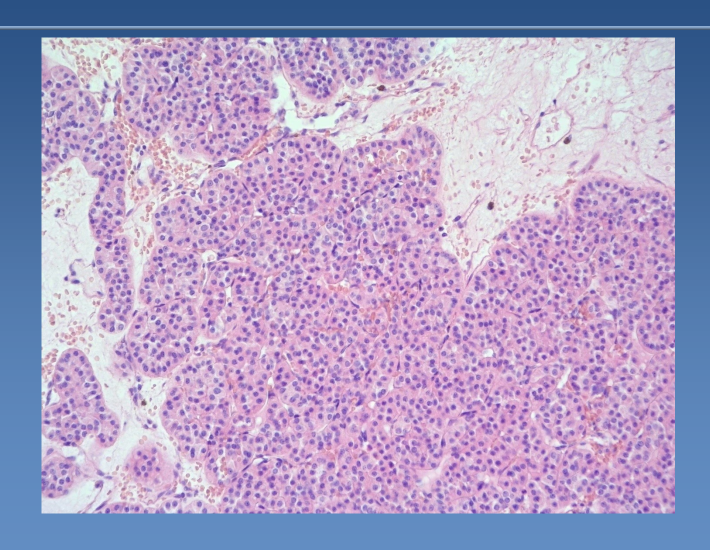


⇒ renal oncocytoma

- •gross: demarcated tumor of red-brown colour, variable size central scar
- micro: eosinophilic, granular cytoplasm, cells in acinar, tubular, solid nests; central hyalinne scar

Renal oncocytoma





Renal cell carcinoma (RCC)



- More common in males; middle-older age
- Smoking as major risk factor
- mostly sporadic tumors, 4% part of hereditary syndromes



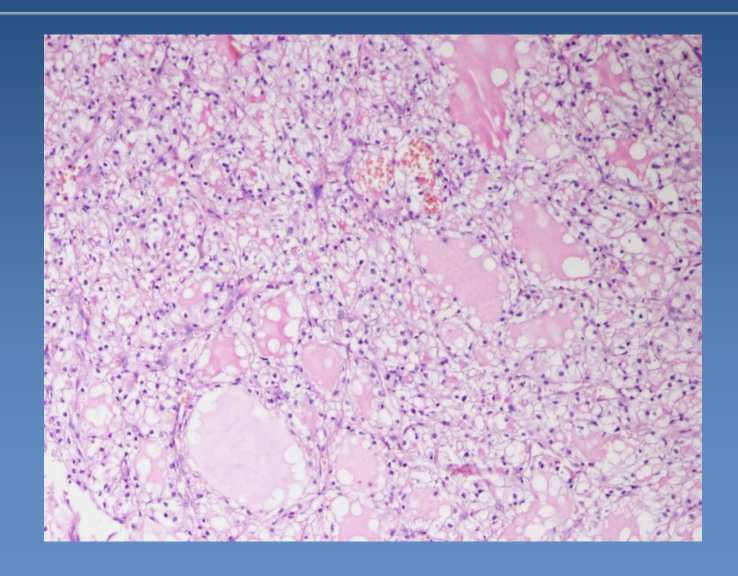
⇒ 70-80% of all RCC

- •gross: demarcated tumor, yellowish colour commonly with haemorrhagic, necrotic, fibrotic foci
- angioinvasive tendency direct grow into renal vein, vena cava;
- invasion into pelvis haematuria



- •Metastases via blood mostly (lungs, bones, brain)
- •micro: large cells with clear granular cytoplasm (glycogen
- + lipids)







- clinical: local symptoms late, haematuria. Fever, paraneoplastic syndromes
- prognosis according to the tumor size/stage
- ca < 3 cm quite good</p>

Papillary RCC

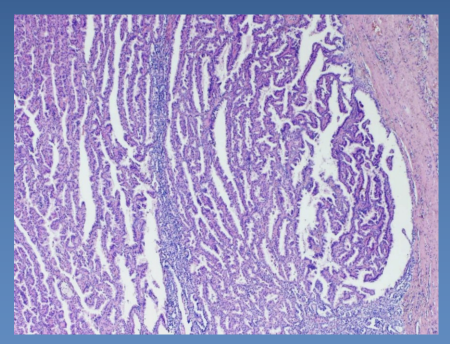


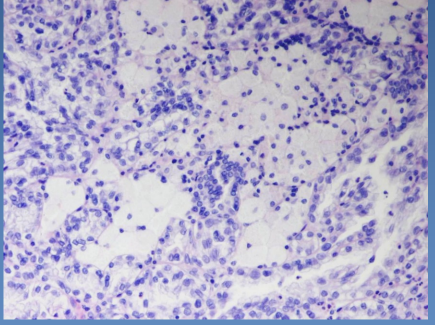
⇒ 15% of all RCC

- •gross: well-demarcated, regressive changes, commonly multifocal and bilateral
- micro: malignant epithelial cells covering stromal papillae,with stromal foam macrophages

Papillary RCC









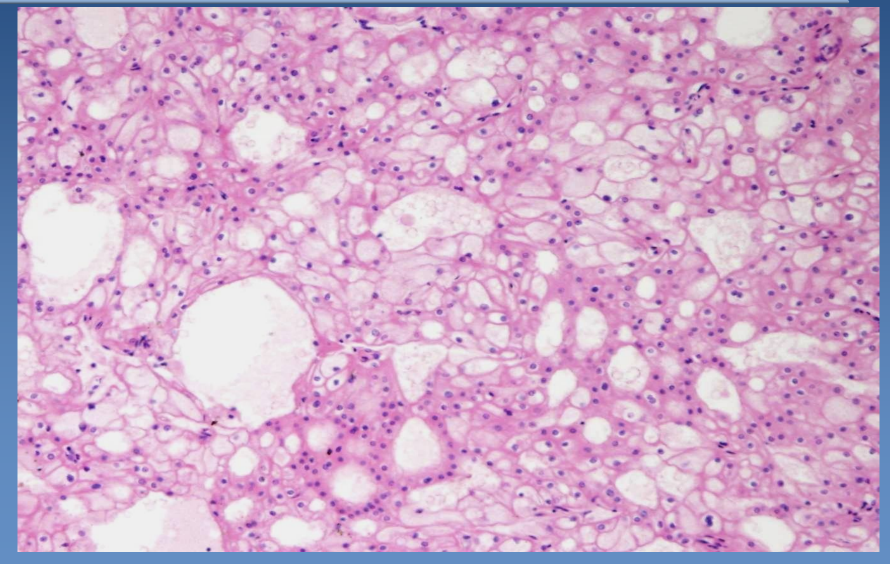


⇒ 5% of RCC.

- •gross: well demarcated, partial lobulisation, brown colour
- micro: eosinophilic granular cytoplasm, distinctive cell membranes, shrunken ("raisin") nucleus

Chromophobe RCC







- 3rd most common malignant pediatric tumor
- Diagnosed mostly in the 3rd-4th year of age
- Sporadic, or part of some syndromes

•gross: large, well demarcated tumor, greyish colour, regressive changes





micro: structures attempting to recapitulate variable stages of nephrogenesis

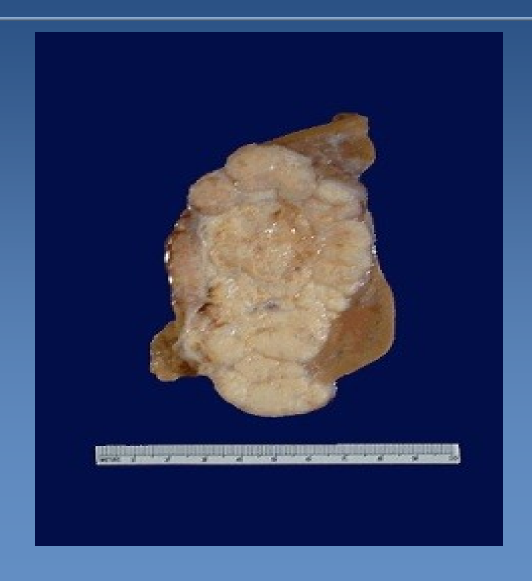
- -Triphasic combination of blastemal, stromal and epithelial cell types in variable percentage
- Highly cellular foci resembling embryonal blastema divided by strands of immature mesenchyme



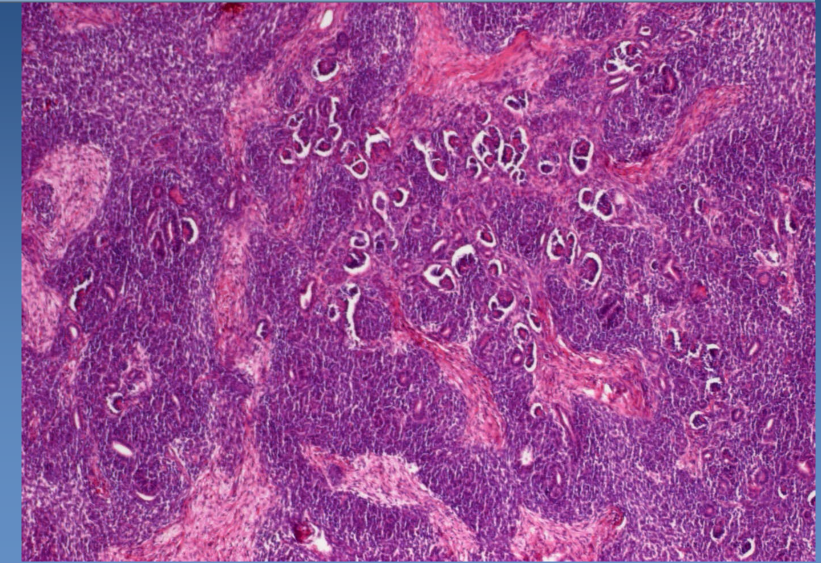


- clinical: large tumor, palpable, complications due to compression of adjacent organs, hematuria
- prognosis: good, CHT (RT carefully, second malignancies possible)

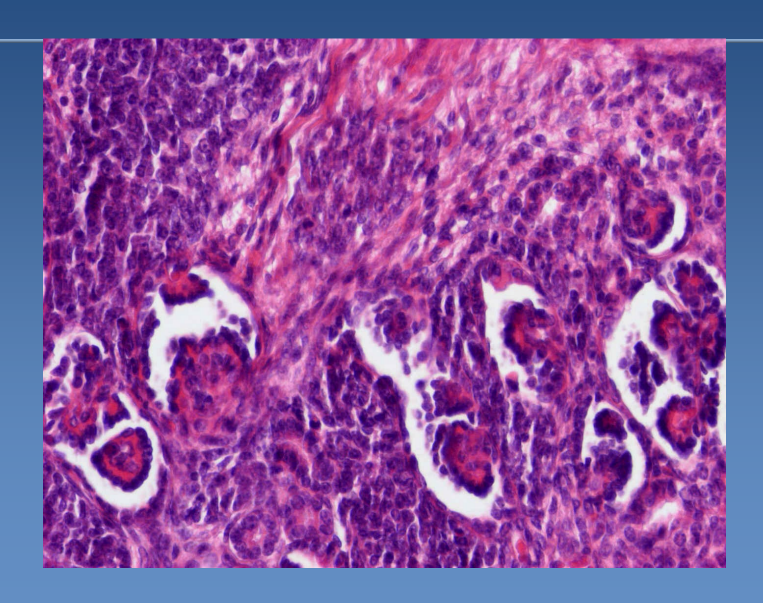












Urinary tract disorders



Urinary tract



- **⇔** Calices
- ⇒ Pelvis
- **⇒** *Ureters*
- ⇒ Urinary bladder
- ⇒ Urethra

Inflammations



- Mostly ascending infection
 - ⇒ urethritis
 - urocystitis
 - possible progression into kidney
 - etiology: E.coli, Proteus, Klebsiella, Enterococcus, Neisseria gonorrhoeae, etc.
 - Candida, Schistosoma,

Inflammations



- dysuria, polakisuria (urging), raised temperature
 - •gross: haematuria, pyuria
 - Hypeaemic mucosa, possible pseudomembrane, ulceration
- complications: progression of inflammation into adjacent structures: glands, interstitium – phlegmona, periurethral abscess

Inflammations



- micro:
 - acute inflammation with prevalence of neutrophils,
 regressive changes of transitional cell epithelium
 - chronic inflammations reactive changes of transitional cell epithelium, squamous/glandular metaplasia. Brunn nests – cystitis cystica
- urethra caruncula urethrae pseudotumorous hyperplastic polyp in the region of urethral orifice.





- Pathological dilatation of the renal pelvis and calyces
 - Causes:
 - Impacted stone, ...
 - -Tumors
 - External compression (pregnancy, prostatic hyperplasia, ...)

Tumors



- * benign x low malignant potential x frankly malignant
- * flat x papillary lesions
 - → Mostly urothelial

Precursor lesions:

- ⇒ Urothelial dysplasia
- risk factors:
 - M:F 3:1
 - smoking
 - professional exposure (aromatic amines, etc.)



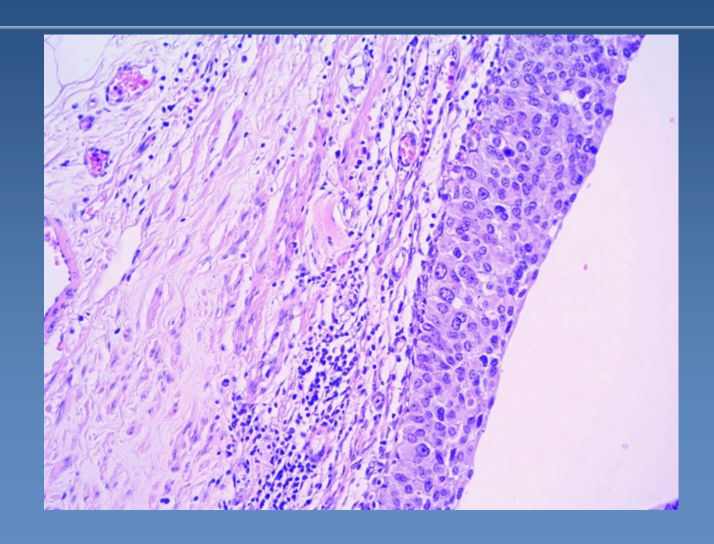


 Micro: flat lesion, cytologic atypia with loss of cell polarity, ↑ mitotic activity in upper layers of urothelium, ↑ N/C ratio, coarse chromatin

*LG (low grade) IUN (intraurothelial neoplasia) x HG IUN (CIS)











urothelial papilloma

Solitary papillary lesion covered by normal urothelium without cytological or architectonic atypias.



Papillary urothelial neoplasm

- ⇒ papillary urothelial neoplasm of low malignant potential (PUNLMP)
 - recurrent tumor
 - papillae covered by hyperplastic urothelium with preserved stratification, minimal cytonuclear atypia, sporadic mitoses.

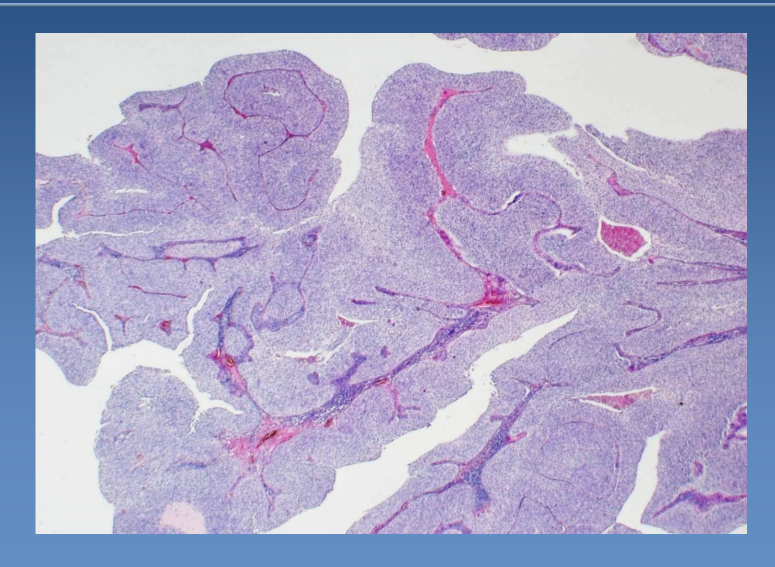
Papillary urothelial neoplasm



- □ non-invasive papillary urothelial carcinoma
 - low grade
 - high grade
- → Papillary neoplasia without signs of invasion into stroma (suburothelial mesenchymal tissue)
- \Rightarrow LG
- altered papillary architectonics,
- mild cytonuclear atypia
- basal layer mitoses

Low grade non-invasive papillary urothelial carcinoma





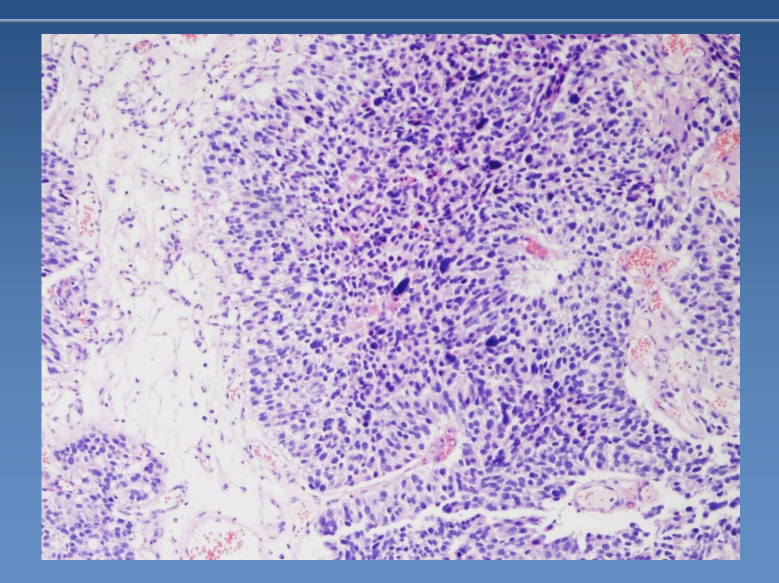
Non-invasive papillary urothelial carcinoma



- \Rightarrow HG
- papillary fusion, solid foci
- loss of cell polarity
- moderate high grade of anisocytosis and anisokaryosis
- atypical mitoses in upper layers of neoplastic epithelium

High grade urothelial carcinoma











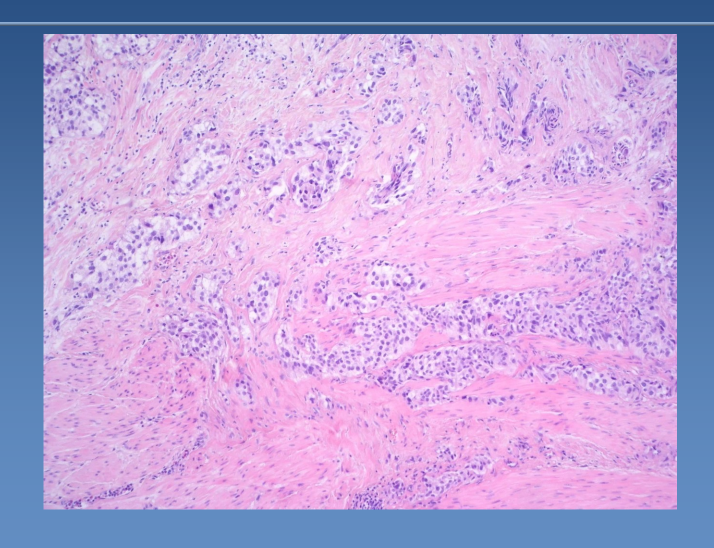
Invasive (infiltrating) urothelial carcinoma



ca invasion into sub-urothelial fibrotic tissue or deeper (muscle, ...)

Invasive urothelial carcinoma









* Less common carcinomas

- ⇒ squamous cell carcinoma (schistosomiasis)
- ⇒ adenocarcinoma
- neuroendocrine carcinoma

Mucinous adenocarcinoma



