# **MEDICINE**

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# Medicine

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			3	Dermatology	60
				3.1 Anatomy and physiology	60
				3.2 Principles of management of	
				skin disease	61
				3.3 Skin cancers	63
(	Contents			3.4 Fungal infections	63
•	Lonients			3.5 Scabies	64
				3.6 Acne	64
				3.7 Eczemas	65
				3.8 Psoriasis	66
C	ontents	1		3.9 Hypopigmentation	69
	ontents	1		3.10 Hyperpigmentation	69
Pı	reface	3		3.11 Stuff not large enough to devote	0,
				a section to	70
1	Respiratory medicine	4		d Section to	70
	1.1 Tuberculosis	4	4	Nephrology	71
				4.1 Anatomy and Physiology	71
2	Cardiology	5		4.2 Presenting problems in urinary	
	2.1 Anatomy and physiology	5		disease	72
	2.2 Investigation of CVS disease	7		4.3 Glomerular diseases ("Glomeru-	
	2.3 Presenting problems in CVS dis-			lonephritides")	73
	ease	8		4.4 Nephrotic syndrome	74
	2.4 ECG	14		4.5 Acute post-streptococcal	
	2.5 Coronary Artery Disease	17		glomerulonephritis	77
	2.6 Arrhythmias	18		4.6 Alport's syndrome	79
	2.7 Atrial fibrillation	21		4.7 Adult Polycystic Kidney Disease	79
	2.8 AV block	22		4.8 Renal artery stenosis	81
	2.9 Antiarrhythmics	23		4.9 Thrombotic microangiopathies	
	2.10 Coronary artery disease	25		(HUS, TTP)	82
	2.11 Heart failure	29		4.10 Acute kidney injury (AKI)	83
	2.12 Peripheral artery disease (PAD) .	33		4.11 Chronic kidney disease (CKD)	85
	2.13 Buerger's and Raynaud's	35		4.12 UTI	90
	2.14 Aortic diseases	36		4.13 Renal cell cancer	91
	2.15 Hypertension	41		4.14 Stuff not large enough to devote	
	2.16 Acute rheumatic fever	41		a section to	93
	2.17 Mitral valve disease	44			
	2.18 Aortic valve disease	47	5	Rheumatology	94
	2.19 Tricuspid valve disease	49		5.1 Investigation of musculoskele-	
	2.20 Pulmonary valve disease	49		tal disease	94
	2.21 Prosthetic valves	50		5.2 Seropositive vs Seronegative	
	2.22 Infective endocarditis	50		arthritis	95
	2.23 Congenital heart disease	52		5.3 Osteoarthritis	95
	2.24 Myocarditis	54		5.4 Spondyloarthropathies	96
	2.25 Cardiomyopathy	56			
	2.26 Cardiac tumours	57	6	Neurology	98
	2.27 Pericardial diseases	57		6.1 Raised ICP	98
				6.2 Neurological emergencies	99

2 CONTENTS

	6.3	Status epilepticus 1	00
	6.4	All jerks root values 1	00
	6.5	Subarachnoid haemorrhage 1	
	6.6	Subacute combined degeneration 1	
	6.7	Cauda equina and Conus	
		medullaris lesions 1	03
	6.8	Neurogenic Bladder 1	05
	6.9	Parkinson's disease 1	
7	Dial	betes Mellitus 1	12
	7.1	Mechanism of insulin secretion . 1	
	7.2	Incretin effect 1	
	7.3	Diagnostic criteria 1	13
	7.4	Diabetic ketoacidosis (DKA) 1	14
	7.5	Hypoglycaemia	
	7.6	Insulin therapy	
	7.7	Oral Hypoglycaemic Agents 1	
	7.8	Complications of DM 1	
	7.9	Pathogenesis of chronic compli-	
		cations	19
	7.10	Stuff not large enough to devote	
		a section to	21
8	Gast	trointestinal diseases 1	22
	8.1	Weight loss	22
9	Hae	matology 1	<b>2</b> 3
	9.1	Chronic myeloid leukaemia	
			<b>2</b> 3
10	Nut	ritional diseases 1	<b>2</b> 5
		Vitamins	25

## **Preface**

What follows are my own notes on various topics in medicine, mostly based off of the following sources:

- ▶ Davidson's Principles and Practice of Medicine 23rd ed.
- ► *Kumar and Clark's Clinical Medicine* 10th ed.
- ► *Oxford Handbook of Clinical Medicine* 10th ed.
- ► Long Cases in Clinical Medicine 2nd ed.
- ► Short Cases in Clinical Medicine 6th ed.

The date below tells you the last time when I edited this document, so refer to that if you're worried about the temporal validity of the contents. The chapters are arranged somewhat at random, partly reflecting the order in which I studied them. These notes are, first and foremost, for my personal use, so pardon the inconvenience. Over time I will try tidying things up more. There's probably plenty of mistakes, all my own. Inform me whenever you find one.

Use at your own peril.

Susmit Islam 2022-09-15

# Chapter 1

# Respiratory medicine

## 1.1 Tuberculosis

## Side effects of anti-TB drugs

- ► Isoniazid:
  - o Hepatitis
  - o Rash
  - $\circ$  B<sub>6</sub> deficiency  $\to$  peripheral neuropathy (so pyridoxine supplement required during therapy)

## ► Rifampicin:

- o Hepatitis
- o Rash

## **▶** Pyrazinamide:

- o Hepatitis
- o Hyperuricaemia (rarely turns into gout)

#### **▶** Ethambutol:

- Retrobulbar neuritis (reversible)
- o Arthralgia

## **▶** Streptomycin:

- Ototoxicity (8th nerve palsy)
- o Rash

# Chapter 2

# Cardiology

## 2.1 Anatomy and physiology

## Coronary circulation

- ▶ RCA  $\rightarrow$  RA, RV, SA node (in 60% individuals), AV node (90%)
  - Posterior descending artery → posteroinferior part of interventricular septum and posterior LV
    - Branch of RCA (in 90%) or LCX (in 10%)
- ▶ LAD → anterior superior part of interventricular septum, anterior wall of LV
  - LAD is called the widowmaker artery, for almost the entirety of LV is supplied by it, and so infarctions involving this have high fatality.
- ▶ LCX  $\rightarrow$  lateral, posterior and inferior LV
- ► See also: table 2.2

## Electrophysiology of the heart

#### Sinoatrial node

- ► Spontaneous depolarization = *pacemaker potential* (details: Fig. 2.1)
  - Due to small influx of Na<sup>+</sup> ions, with 2 components
  - $\circ$  Background inward current  $I_b$
  - o "Funny" current  $I_f$ 
    - "Funny" as the channels are activated in hyperpolarized cells, as opposed to most other voltage-gated channels, which activate upon depolarization
- ▶ Atrial tissue is activated like a "forest fire", but by the time the potential reaches the insulating annulus fibrosus at the AV junction, it peters out (which is why we need the AV node to act as a "repeater")

## Action potentials in the conductive system and the myocytes



Figure 2.1: Cardiac action potentials. CaL: L-type (long-lasting)  $Ca^{2+}$  channel, CaT: T-type (transient)  $Ca^{2+}$  channel.  $I_{K1}$ : inward rectifier  $K^+$  current,  $I_{Na}$ : inward  $Na^+$  current,  $I_{To}$ : transient outward  $K^+$  current,  $I_{CaL}$ : inward  $Ca^{2+}$  current,  $I_K$ : delayed rectifier (outward)  $K^+$  current.

## Effects of the autonomic nervous system on the heart

- $ightharpoonup eta_1$ -adrenergic stimulation  $ightharpoonup \uparrow$  force of contraction, heart rate (+ve inotropic and chronotropic)
  - o  $\beta_1$  stimulation  $\to \oplus$  adenlylyl cyclase-cAMP system  $\to \oplus$  intracellular protein kinases  $\to \uparrow$  phosphorylation of proteins including L-type Ca<sup>2+</sup> channels  $\to$  **enhanced** Ca<sup>2+</sup> **influx**  $\to \uparrow$  FoC
  - o Return of Ca<sup>2+</sup> to sarcoplasmic reticulum from the myocyte is mediated by phospholamban, which enhances Ca<sup>2+</sup> reuptake into the SR in its phosphorylated state. So  $\beta_1$  stimulation also promotes Ca<sup>2+</sup> removal from the myocyte, thereby enhancing myocardial relaxation. Enhanced relaxation  $\rightarrow \uparrow \uparrow$  ventricular filling, and the  $\uparrow$ FoC helps the heart to pump that extra blood out properly.

## Cardiac peptides

- ► **ANP** (atrial natriuretic peptide)
  - Released by atrial myocytes upon being stretched
  - $\circ$  Vasodilator  $\rightarrow \downarrow BP$
  - o Diuretic (↑Na<sup>+</sup> and H<sub>2</sub>O excretion)
- ► **BNP** (brain/B-type natriuretic peptide)
  - o Called so because first discovered in porcine brain extracts
  - o Released by ventricular myocytes upon being stretched
  - Diuretic
- ► Neprilysin:
  - o NOT a cardiac peptide (made by the kidney and other tissues)
  - o Breaks down ANP, BNP  $\rightarrow$  vasoconstrictor
  - o Therapeutic target in resistant heart failure

## Haemodynamic effects of respiration

#### **Effects**

	Inspiration	Expiration
JVP	<b>+</b>	<b>†</b>
BP	↓ (upto 10 mmHg)	<b>†</b>
HR	<b>†</b>	<b>↓</b>
2nd heart sound (S <sub>2</sub> ) splitting	<b>†</b>	<b>↓</b>
Right sided murmurs	<b>†</b>	<b>↓</b>
Left sided murmurs	<b>↓</b>	<b>†</b>

Table 2.1: Haemodynamic effects of respiration

#### Mechanisms

- ▶ As the circulatory system is a closed system, if any compartment within it has more blood at any given time, the other compartments will have less blood.
- ► BP:
  - $\circ$  Inspiration  $\to$  -ve intrathoracic pressure  $\to \uparrow$  venous return to RA  $\to \uparrow$  RV output
  - o Inspiration  $\to \uparrow$  pulmonary vascular capacitance  $\to \downarrow$  venous return to LA  $\to \downarrow$  LV output  $\to \downarrow$  BP
- ▶  $S_2$  splitting (i.e. delay between  $A_2$  and  $P_2$ ):
  - o Inspiration  $\to \uparrow$  RV filling  $\to \uparrow$  prolonged ejection from RV  $\to \uparrow$  delayed  $P_2$
  - o Inspiration  $\to \ \downarrow$  LV filling  $\to \ \downarrow$  faster ejection from LV  $\to \ \downarrow$  faster  $A_2$

## 2.2 Investigation of CVS disease

#### **ECG**

▶ Discussed in Sec. 2.4

## Chest X-ray<sup>1</sup>

- ► Cardiomegaly: if cardiothoracic ratio (CTR) > 0.5
- ► Findings according to the chamber enlarged:
  - Left atrial enlargement:
    - Straight left heart border

<sup>&</sup>lt;sup>1</sup>A very nice intro to CXR interpretation from Axis Medical School. This playlist by Medzcool is also really good.

- Double cardiac shadow to the right of sternum
- Widened carinal angle
- o Right atrial enlargement:
  - Projects from right heart border to right lower lung field
- o Left ventricular enlargement: may be due to dilation or hypertrophy
  - Cardiomegaly
  - Rounding of left heart border if hypertrophy
- o Right ventricular enlargement: dilation or hypertrophy
  - Cardiomegaly
  - Upwards displacement of apex
  - Straight left heart border

## 2.3 Presenting problems in CVS disease

#### **Pulse**

- ► Assess
  - o Rate
  - o Rhythm
  - o Volume: best assessed in the carotids
  - o Character: best assessed in the carotids
  - o Condition of the vessel wall
  - o Radio-radial and radio-femoral delays
- ▶ Volume
  - o High volume pulse
    - AR
    - Hyperdynamic circulation
  - o Low volume pulse
    - Shock
    - AS, MS
    - Constrictive pericarditis
    - Cardiac tamponade
    - Pulmonary htn
- ► Character
  - o Anacrotic:
    - Slow rising, low volume
    - Found in AS
  - o Dicrotic:
    - Double peak, one systolic, other diastolic
    - Found in enteric fever
  - o Bisferiens:

- Double peak, both systolic
- Found in combined AS and AR
- o Waterhammer / Collapsing:
  - Rapid rise followed by rapid fall
  - Accentuated by raising the arm
  - Seen in AR
- o Paradoxus:
  - Cardiac tamponade
  - Chronic constrictive pericarditis
  - Acute severe asthma and COPD
  - Massive PE

## Heart sounds<sup>2</sup>

 $S_1$ 

- ► Cause: closure of atrioventricular valves (mitral and tricuspid)
- ► Timing: onset of ventricular systole
- ► Best heard at: the *apex*
- **▶** Abnormalities:
  - o Loud: MS (mitral stenosis)

 $S_2$ 

- ► Cause: closure of the semilunar valves (aortic and pulmonary)
- ► Timing: end of ventricular systole
- ► Best heard at: *left sternal edge*
- ► Normally louder and higher-pitched than S<sub>1</sub>
- ightharpoonup A<sub>2</sub> louder than P<sub>2</sub> (details in this subsection)
- ► Abnormalities:
  - Quiet: AR (aortic regurgitation)
  - o Loud: systemic / pulmonary htn

## Splitting of S<sub>2</sub>

- Normally splits as  $S_2 = A_2 + P_2$ , because the left ventricle contracts more forcefully and so the aortic valve closes earlier than the pulmonic valve ("physiological splitting")
- ▶ During inspiration, the right heart receives more blood and the left heart receives less blood, and so  $A_2$  happens even earlier and  $P_2$  even later, leading to increased split.

 $S_3$ 

► Cause: *rapid ventricular filling* immediately after opening of AV valves

<sup>&</sup>lt;sup>2</sup>This video by Dirty medicine discusses heart sounds and murmurs together with detailed explanations. This playlist by Medzcool contains audio recordings of normal heart sounds and the most important abnormal heart sounds and associated murmurs.

- ► Timing: early diastolic
- ▶ Best heard at:
  - o bell at apex
  - o bell because it's low-pitched
- ► Heard as "lub-dub-dum"
- ► Normal in:
  - o Children
  - Young adults
  - o Febrile patients
  - o Pregnancy
- $\blacktriangleright$  **Abnormalities:** presence of S<sub>3</sub> usually pathological after 40y age
  - o LVF
  - o MR (mitral regurgitation)

#### $S_4$

- ► Cause: forceful atrial contraction against non-compliant / stiff ventricle
- ► Timing: *end-diastolic* (just before S<sub>1</sub>)
- ▶ Best heard at:
  - o bell at apex
  - o bell because it's low-pitched
- ► Always pathological
  - o LVH (due to htn, aortic stenosis, or HCM / hypertrophic cardiomyopathy)

#### Added sounds

- ► Opening snap:
  - $\circ$  early diastolic, just after  $S_2$
  - o in mitral stenosis
  - o diaphragm at apex
- ► Midsystolic clicks:
  - o in mitral valve prolapse
  - o diaphragm at apex
- ► Pericardial / friction rub:
  - o in acute pericarditis
  - o diaphragm at any part of precordium with breath held at expiration

## Murmur<sup>3</sup>

#### **Basics**

- ▶ Points to assess: SCRIPT
  - o Site: aortic, pulmonary, tricuspid or mitral area
  - o Character: soft / blowing / decrescendo / crescendo-decrescendo
  - o Radiation: MR  $\rightarrow$  axilla, AS  $\rightarrow$  neck (carotid)
  - o Intensity / grading:
    - 1: very soft (audible in ideal conditions)
    - 2: soft
    - 3: moderate
    - 4: loud, with thrill
    - 5: very loud
    - 6: heard without stetho
  - o Pitch: low (caused by low velocity blood flow) / high (high velocity blood flow)
  - o Timing: systolic / diastolic
- ▶ Right heart murmurs → louder during inspiration (**RINspiration**)
- ▶ Left heart murmurs → louder during expiration (**LEXpiration**)

#### Features of benign murmur

- ► Soft
- ► Midsystolic
- ► Heard at left sternal edge
- ► No radiation
- ▶ No other cardiac abnormalities

#### **Systolic murmurs**

- **▶** Ejection systolic murmurs
  - o Causes: ventricular outflow obstruction
    - AS (aortic stenosis)
    - PS (pulmonary stenosis)
  - Occur during mid-systole with crescendo-decrescendo pattern (i.e. gradual rise followed by gradual fall)
- **▶** Pansystolic murmurs
  - o Causes: leakage of blood into low pressure chamber from a ventricle
    - MR (mitral regurgitation)
    - TR (tricuspid regurgitation)
    - VSD (ventricular septal defect)

<sup>&</sup>lt;sup>3</sup>Here's some awesome reviews by ZeroToFinals and DirtyMedicine on murmurs. This one by Strong Medicine covers the topic at a slightly deeper level with 5/6 exercises with actual recordings of murmurs.

#### Diastolic murmurs

- ▶ Low-pitched, often difficult to hear so should be examined with the bell
- ► Mid-diastolic murmur
  - o Causes:
    - MS (mitral stenosis): at the mitral area and radiated to the axilla
    - TS (tricuspid stenosis): at the left sternal edge

## ► Early diastolic murmur

- o Soft, blowing, decrescendo pattern (gradual fall)
- o Causes:
  - AR (aortic regurgitation)
  - PR (pulmonary regurgitation)
- o In case of "systolic regurgitations" i.e. MR/TR, due to the high flow throughout systole, the murmur is pansystolic. In case of "diastolic regurgitations", the blood has already mostly flown out of the aorta or the pumonary trunk, leaving little blood for backflow. So in this case, it's early diastolic.

#### **Continuous murmurs**

► Cause: PDA (persistent ductus arteriosus)

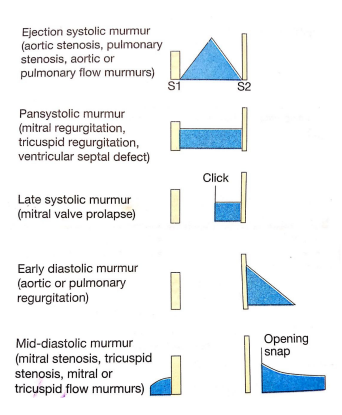


Figure 2.2: The timing and pattern of heart murmurs

## JVP<sup>4</sup>

- ► Reflects rt atrial pressure
- ► Parts of the waveform: fig. 2.3
  - o a wave: Atrial systole
    - Giant "cannon" a wave: CHB (AV dissociation → sometimes atrial and ventricular contraction coincide → rt atrium contracts against closed tricuspid → ↑↑ pressure → giant a wave)
    - Large a wave: pulmonary htn, PS
    - Absent a wave: AF
  - o **c wave:** Atrial relaxation coinciding with ventricular systole (before closure of the tricuspid valve)
  - **x descent:** Atrial relaxation coinciding with ventricular systole (after closure of the tricuspid valve)
  - o **v wave:** Atrial filling
    - Large v wave: TR
  - o **y descent:** Atrial emptying
    - During atrial systole, two opposing forces are at play. Atrial contraction raises the JVP, while blood leaving the compartment lowers it. During the initial part of systole, the contraction is weaker, so the emptying effect takes the lead, giving rise to the *y descent*. Later, the contraction effect is stronger than the emptying effect, and so there's a peak (*a wave*).
- ▶ **Hepatojugular reflux**: pressing on the liver (rt hypochondrium) → ↑emptying of veins into the rt atrium → transient ↑rt atrial pressure
  - If RVF, then the increase in JVP is more sustained

## **▶** JVP vs carotid pulse

- JVP biphasic (2 peaks within each cardiac cycle, one during atrial systole, other during atrial diastole), carotid monophasic (peak only during ventricular systole)
- o JVP impalpable but occludable, carotid palpable but non-occludable
- o JVP height varies with respiration, carotid pulse doesn't.

<sup>&</sup>lt;sup>4</sup>This video by ZeroToFinals explains this topic far better than anywhere I've read or seen.

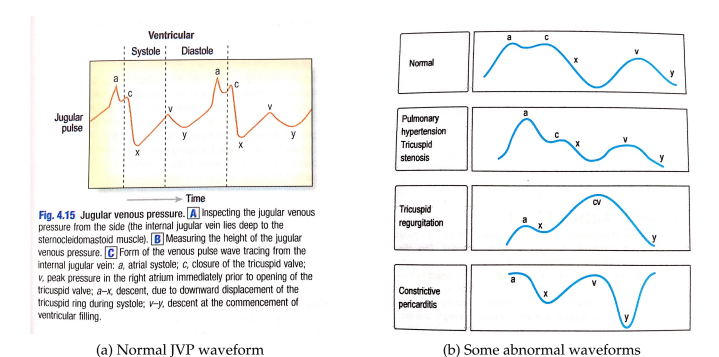


Figure 2.3: JVP

## 2.4 ECG

## Anatomy of an ECG

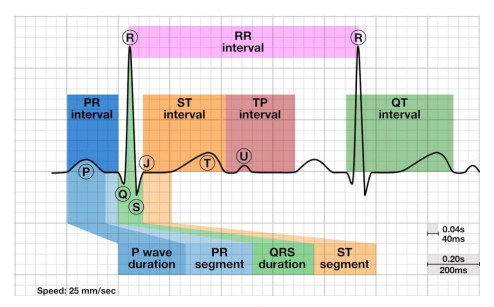


Figure 2.4: Parts of a normal ECG

2.4. ECG 15

## Abnormalities of components

## Pathological Q

- ► Depth > 2mm
- ► Height > 1mm
- ▶ Present in  $\geq$  2 leads
- ▶ Assocd with loss of R height (Q > R/4; normally  $Q \le R/4$ )
- ► Indicates transmural myocardial necrosis

## Segments vs intervals

- ▶ e.g. ST segment = end of  $S \rightarrow start$  of T
- ▶ PR interval = start of P  $\rightarrow$  start of R

#### ST segment elevation

- ▶ Normal: upto 1mm in limb leads, upto 2mm in chest leads
- **▶** Causes
  - **STEMI:** convexity upwards
  - o Acute periCArditis:: conCAvity upwards
- ► Indicates ongoing myocardial injury

## Myocardial infarction

A somewhat interesting physiological explanation on how the changes arise

#### Sites of infarction based on lead

Septal: V<sub>1</sub>, V<sub>2</sub>
 Anterior: V<sub>3</sub>, V<sub>4</sub>

Lateral: I, aVL, V<sub>5</sub>, V<sub>6</sub>
Extensive anterior: V<sub>1</sub>-V<sub>6</sub>
Anterolateral: I, aVL, V<sub>1</sub>-V<sub>6</sub>

► Further details: table 2.2 and subsection 2.1

Location	Leads	Coronary artery
Anterior	$V_2$ - $V_4$	LAD
Lateral	$V_5$ - $V_6$	LCX > LAD
Inferior	II, III, aVF	RCA ≫ LCX
Posterior	V <sub>1</sub> , V <sub>2</sub> (depression)	$RCA \gg LCX$

Table 2.2: ECG leads and arteries involved according to anatomical location of infarct

## Reciprocal changes

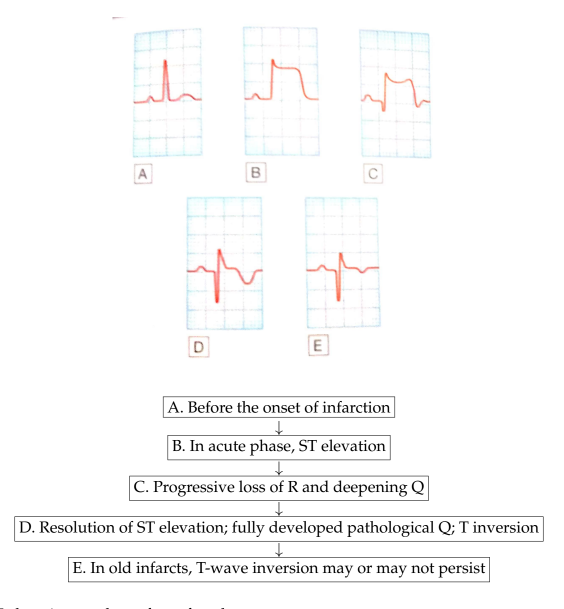
 $\blacktriangleright$  Acute STEMI in some surface of the heart  $\rightarrow$  ST elevation in corresponding leads, and ST depression in reciprocal leads

Site	Facing	Reciprocal
Septal	$V_1, V_2$	V <sub>7</sub> -V <sub>9</sub>
Anterior	$V_3$ , $V_4$	None
Lateral	$I$ , $aVL$ , $V_5$ , $V_6$	II, III, aVF
Inferior	II, III, aVF	I, aVF
Posterior	$V_7$ - $V_9$	$V_1, V_2$

Table 2.3: Reciprocal changes according to location of infarct

## Evolution over time of the ECG appearance of STEMI

► STEMI = complete proximal occlusion of major coronary artery



► ST elevation resolves after a few days

#### **NSTEMI**

- ▶ Partial occlusion of major or complete occlusion of minor coronary artery
- ightharpoonup Subendocardial/partial-thickness MI ightharpoonup no pathological Q
- ► ST depression + T inversion in chest leads

## 2.5 Coronary Artery Disease

▶ Diseases arising due to narrowing of the lumen of one or more coronary arteries and the resulting ischaemia/infarction of the myocardium or the conductive system.

## ► Types:

- o Stable angina: Fixed atheromatous stenosis
- o Unstable angina:
  - dynamic obstruction
  - due to plaque rupture/erosion with thrombosis
- o MI
- Heart failure
- o Arrhythmia
- Sudden cardiac death
  - ventricular arrhythmia
  - asystole
  - massive MI

## 2.6 Arrhythmias

## **Antiarrhythmics**

Discussed in Section 2.9

## Sinus arrhythmia

- ▶ Physiological alteration of HR with respiration
- ▶ Inspiration  $\rightarrow \uparrow HR$ , expiration  $\rightarrow \downarrow HR$  (see also: Table 2.1)
- ► Absent in diabetic neuropathy

## Classification according to ECG morphology

- ► Narrow complex: QRS < 120ms (3 small sqs)
  - o Sinus tachycardia
  - o Atrial fibrillation (irregular narrow complex tachycardia)
  - o Atrial flutter
  - o AV Nodal Re-entry Tachycardia (AVNRT aka SVT)
- ► **Broad complex**: QRS > 120ms (3 small sqs)
  - Ventricular tachycardia
  - o AV Re-entry Tachycardia (AVRT e.g. Wolff-Parkinson-White syndrome)
    - Abnormal band of conductive tissue connecting atria and ventricles (accessory pathway)

## SVT (AVNRT)

## Management

- ► Carotid sinus massage or
- ▶ Valsalva manoeuvre

2.6. ARRHYTHMIAS

- ▶ If the manoeuvre fails,
  - o Adenosine (3-12mg IV) (see more in this section) or
  - o Rate-limiting CCB (Verapamil 5mg IV) or
  - o  $\beta$ -blocker
- ▶ If haemodynamic state compromised, **DC cardioversion**
- ► Recurrent SVT → catheter ablation

#### Atrial flutter

#### Intro

- ► Caused by **macro re-entry circuit** usually in the rt atrium
- ► Atrial rate  $\approx 300$ /min
- ▶ Because of the refractory period, not all of these 300 APs can excite the AV node → 2nd degree type II AV block (2:1, 3:1, or 4:1, corresponding to a ventricular rate of 150, 100, or 75/min)
- ► Saw-tooth flutter waves in ECG

## Management

- $\blacktriangleright$  Medical: digoxin,  $\beta$ -blockers, or verapamil
- ▶ DC cardioversion
- $\triangleright$   $\beta$ -blockers or amiodarone to prevent recurrence
  - o Class Ic antiarrhythmics (e.g. flecainide) contraindicated
    - Risk of facilitating 1:1 AV nodal conduction leading to extreme tachy and haemodynamic compromise
- ► Catheter ablation
- ► Anticoagulant therapy (similar to A-fib's anticoagulant regimen)

## Ventricular tachycardia / VT

- ▶ Most commonly in acute MI, chronic CAD, cardiomyopathy
- ► Caused by abnormal automaticity in ischaemic tissue
- ► Features:
  - Palpitation
  - o Dyspnoea
  - Lightheadedness
  - o Syncope
  - o Hypotension
  - o Cardiac arrest
- ▶ Management: Prompt restoration of sinus rhythm by
- ► If systolic BP < 90 mmHg, **DC cardioversion**

- ightharpoonup Else, IV  $\beta$ -blocker / amiodarone
  - o If medical therapy fails, DC cardioversion

## **Torsades de Pointes**

- ► Form of polymorphic VT arising as a complication of prolonged QT- interval (i.e. prolonged ventricular repolarisation)
- ► Literally, "twisting of the peaks" (French)
- ▶ QRS complexes "twist" around the isoelectric line



Figure 2.5: Torsades de pointes

- ▶ Usually non-sustained, repetitive
- ► May progress to V-fib
- ► Causes of long QT (> 0.44s in men, > 0.46s in women)
  - o Electrolyte imbalance
    - Нуро-К
    - Hypo-Ca
    - Hypo-Mg
  - o Drugs
    - Class Ia, Ic antiarrhythmics (e.g. disopyramide, flecainide etc)
    - Class III antiarrhythmics (amiodarone, sotalol)
  - o Congenital long QT syndrome
- ► Management: IV Mg

## 2.7 Atrial fibrillation

#### Intro

- ► Most common sustained arrhythmia
- ► Associated with underlying heart disease, systemic embolism, stroke

#### **Causes**

- ► Cardiac
  - o CAD (including acute MI)
  - o Mitral stenosis (MS; rheumatic mitral valve disease)
  - Hypertension
  - o Cardiomyopathy
- ► Non-cardiac
  - o Thyrotoxicosis
  - o Pulmonary embolism
  - o Pneumonia
  - o Alcoholism

## Investigation

- ► ECG
- ► Echo: to see valvular condition
- ► Thyroid function test: to exclude thyrotoxicosis

## Management of AF

- ► Rhythm control:
  - o Pharmacological cardioversion
    - Pt stable + no history of heart disease  $\rightarrow$  **IV flecainide**
    - Structural / ischaemic heart disease → IV amiodarone
  - o DC cardioversion / catheter ablation if drugs fail
- ▶ Rate control:
  - o Digoxin
  - $\circ$   $\beta$ -blockers
  - Rate-limiting CCB: verapamil / diltiazem
- **►** Thromboprophylaxis:
  - o Oral Warfarin
  - o Target INR: 2.0-3.0
  - o Start 4wks before cardioversion, continue till 3mo after successful cardioversion
  - o Monitor stroke risk by CHA<sub>2</sub>DS<sub>2</sub>-VASc score, and bleeding risk by HAS-BLED score (HAS-BLED  $\geq$  3  $\rightarrow$  close monitoring on anticoagulant)

- o Directly acting oral anticoagulants (DOACs):
  - Factor Xa inhibitors: Rivaroxaban, apixaban, edoxaban
  - Direct thrombin inhibitors: dabigatran
  - Efficacy ≥ warfarin, haemorrhage risk, drug interactions ≪ warfarin, effective antidote to manage acute bleed (idarucizumab)
- o Avoid aspirin: little effect on embolic stroke, significant bleeding risk

## 2.8 AV block

## 1° AV block

- ► Prolonged PR interval (> 0.2s)
- ► Rarely symptoms; rarely need treatment

#### 2° AV block

- ▶ **Dropped beats** i.e. not all impulses in the atria are conducted to the ventricles
- ► Mobitz I:
  - o Progressively prolonging PR, then dropped beat, repeat
  - o AKA Wenckebach phenomenon
  - o Caused by disease in the AV node

#### ► Mobitz II:

- o PR constant (when conducted), fixed proportion of atrial impulses conduct to the ventricles (e.g. 3:1 block means for every 3 impulses coming from the atria 1 excites the ventricles)
- o Caused by disease in the His-Purkinje system
- Risk of asystole (cardiac arrest)
- ▶ Impossible to distinguish type I and II in 2:1 block

#### 3° AV block

- ► AV dissociation
- ▶ Causes:
  - o MI
  - o Infective endocarditis
  - o Drugs: Digoxin, β-blockers, CCBs

#### **Features**

- ► Recurrent syncope ("Stokes-Adams" attacks)
- ▶ Brief anoxic seizures

## Management

- ▶ Acute inferior MI often accompanied by AV block, as RCA supplies both the AV node and the inferior wall. Treat MI, no other treatment for AV block required.
- ► Symptomatic 2° and 3° block → atropine
- Asymptomatic 1° and 2° Mobitz I → no treatment required
- ▶ Mobitz II /  $3^{\circ}$  → permanent pacemaker (risk of sudden death from asystole)

## 2.9 Antiarrhythmics

#### Classification

#### By effect on the action potential

- ► Class I: Na<sup>+</sup>-channel blockers (manipulate length of AP by controlling the refractory period)
  - o Ia. Prolong AP: quinidine, disopyramide
  - o Ib. Shorten AP: lidocaine, mexiletine
  - o Ic. No effect on AP: Flecainide, propafenone
    - avoid if previous MI († risk of arrhythmia)
    - flecainide used in AF for rhythm control
      - may cause slow atrial flutter with rapid ventricular rate, so prescribed with an AV-node blocking drug e.g.  $\beta$ -blocker
- ightharpoonup Class II:  $\beta$ -blockers
  - reduce SA node firing rate
  - o atenolol, metoprolol, bisoprolol
  - o Uses:
    - Rate control in flutter and AF
    - Prevent recurrence of VT, SVT
- ► Class III:
  - o amiodarone, dronedarone, sotalol
    - sotalol is a racemic mixture of l-sotalol (nonselective  $\beta$  blocker) and d-sotalol (class III)
    - amiodarone also has I, II, IV activity
    - most effective drug in paroxysmal AF
  - prolong plateau phase → ↑refractory period
  - o prolong QT-interval, may predispose to torsades de pointes / VT
- ► Class IV: rate-limiting CCBs
  - o Verapamil, diltiazem
  - o Uses: prevent SVT, rate control in AF

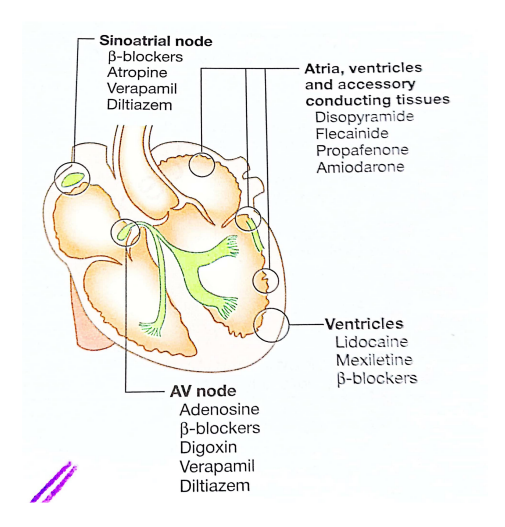


Figure 2.6: Classification of antiarrhythmics by site of action

## By site of action

## Other antiarrhythmics

- ► Atropine: in symptomatic / severe bradycardia
- ► Adenosine:
  - Mechanism: bind to A1 receptors in the AV node producing a transient AV block for a few seconds, which terminates the cyclical propagation of the impulse through the accessory pathway
  - $\circ$  May cause bronchospasm  $\rightarrow$  avoided in asthma
- ► Digoxin:
  - o Slows conduction and prolongs AV node refractory period
    - So used for rate control in AF
  - $\circ$  Enhances conduction and shortens refractory period in other parts of the heart  $\rightarrow$  may promote atrial / ventricular ectopics

## 2.10 Coronary artery disease

## Risk factors

- ► Smoking: most important modifiablle risk factor
- ► Age and sex:
  - o Premenopausal women  $\rightarrow \downarrow$ risk than men
  - o Postmenopausal women  $\rightarrow$  = risk as men
  - HRT in postmenopause ↑risk
- ▶ Diet:
  - $\circ \downarrow$ fruit, veg, PUFA  $\rightarrow \uparrow$ risk

## Angina pectoris

#### Intro

► Chest pain due to transient myocardial ischaemia resulting from an imbalance between myocardial O<sub>2</sub> supply and demand

## **Pathogenesis**

- ► Causes
  - Atherosclerosis
  - Coronary artery spasm
  - o Syndrome X:
    - Effort angina + evidence of ischaemia on stress test + normal arteries on angio
    - Mechanism unknown

#### Investigation

- ► Exercise ECG: first line
  - o Planar / downsloping ST depression of ≥ 1mm indicates ischaemia
  - o Upsloping ST depression nonspecific; may be nor -mal
- ▶ Myocardial perfusion scan / stress echo: if exercise ECG inconclusive
- ► CT angiogram

#### Management

- **▶** Lifestyle changes:
  - Smoking cessation
  - Wt reduction
  - o Regular exercise (up to, but not beyond, point of chest discomfort)
  - o Sublingual GTN before exertion that may induce angina
- ► Pharmacological therapy:

- Start with sublingual GTN +  $\beta$ -blocker
- Add CCB or long-acting nitrate if needed
- Statin, even if lipid profile normal
- o If 2° to CAD, antiplatelet therapy
  - Low-dose aspirin (75mg) / Clopidogrel (75mg) continued indefinitely
- Sublingual GTN:
  - Relieve attack within 2-3 mins
  - Side effects:
    - Headache
    - Hypotension
    - Syncope
  - Continuous therapy may cause tolerance
    - Avoided by 6-8h nitrate free period, best achieved at night as the patient is inactive
- o Beta-blockers:
  - Nonselective may aggravate vasospasm by blocking coronary arterial  $\beta_2$
  - So once daily cardioselective  $\beta$  blocker (metoprolol / bisoprolol)
  - Should not be withdrawn rapidly:  $\beta$ -blocker withdrawal syndrome
    - May precipitate arrhythmias, worsen angina, MI
- o CCBs:
  - ullet Dihydropyridines (e.g. amlodipine) may cause reflex tach so combo with eta-blocker
  - Verapamil and diltiazem can be used as monotherapy as don't cause tach
  - CCB  $\rightarrow \downarrow$  contractility  $\rightarrow$  used with care if poor LV function
- K-channel activators:
  - Nicorandil: only approved drug in this class
    - Vasodilator
    - No tolerance as with nitrates
- o I<sub>f</sub>-channel antagonists:
  - Ivabradine:
    - Induces bradycardia
    - Doesn't inhibit contractility
    - Safe in heart failure

## ► Nonpharmacological therapy

- PCI with / without stenting: if single or dual vessel disease
  - Restenosis is the main complication (in upto 1/3rd cases, usually occurs within 3mo)
  - Prevented by stenting
  - Drug-eluting stents reduce risk further (antiproliferative drugs e.g. sirolimus, paclitaxel prevent neo intimal hyperplasia)
- o CABG: if triple vessel disease

## Acute coronary syndrome (ACS)

#### Intro

► ACS = unstable angina or MI

## Acute management

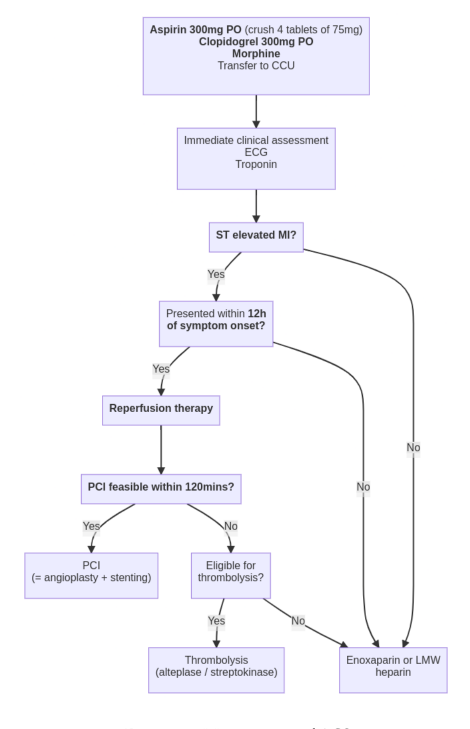


Figure 2.7: Management of ACS

## Long-term management

- ► Lifestyle changes: see here
- **▶** 2° prevention
  - o Antiplatelet therapy: aspirin and/or clopidogrel
  - o ]beta-blocker
  - o ACEi / ARB
  - o Statin
  - o DM control
- ▶ If high-risk, ICD (implantable cardiac defibrillator)

#### **Complications**

## ► Arrhythmias:

- o Usually transient, no prognostic significance
- o VF is the major cause of death in those who die before receiving medical care
- Common arrhythmias in ACS
  - VF
  - V-tach
  - AF
  - A-tach
  - Accelerated idioventricular rhythm
  - Ventricular ectopics
  - Sinus bradycardia
  - AV block
- ► Recurrent angina
- ► Acute heart failure
- **▶** Pericarditis
- ▶ Dressler's syndrome
  - o Fever, pericarditis, pleurisy
  - o Probably autoimmune
  - If prolonged or severe, high dose aspirin, NSAIDs or glucocorticoids

#### ► Papillary muscle rupture

- Causes sudden severe MR, leading to the features
  - Acute pulmonary oedema
  - Shock
  - Pansystolic murmur radiating to axilla
- Confirmed by echo
- Emergency valve replacement

## **▶** Ventricular septum rupture

- o Presents with
  - Sudden haemodynamic deterioration

- Pansystolic murmur: loud, radiating to rt sternal border
- Acute RHF (due to lt to rt shunt)
- o Confirmed by Doppler echo and rt heart catheterisation
- o Emergency surgical repair

## **▶** Ventricular rupture

- o Usually fatal
- **►** Embolism
- ► Ventricular remodelling
  - o Thinning and expansion of infarct  $\rightarrow \uparrow$  stress on remaining ventricle  $\rightarrow$  dilatation and hypertrophy of remaining ventricle
  - ACEi and aldosterone antagonists reduce remodelling
- ► Ventricular aneurysm

## 2.11 Heart failure

## **Pathophysiology**

- ► Hazards of prolonged sympathetic stimulation
  - Cardiomyocyte apoptosis
  - Hypertrophy
  - o Focal myocardial necrosis
  - o Predispose to arrhythmias

## **Causes**

- **▶** ↓ contractility
  - o MI
  - Dilated cardiomyopathy
  - Myocarditis

#### **▶** Outflow obstruction

- o Systemic htn, pulmonary htn
- o Aortic stenosis, pulmonary stenosis
- **▶** Inflow obstruction
  - Mitral stenosis, tricuspid stenosis
- ▶ Volume overload
  - Aortic / mitral regurgitation
  - o VSD
- ► Arrhythmia
  - o A-fib

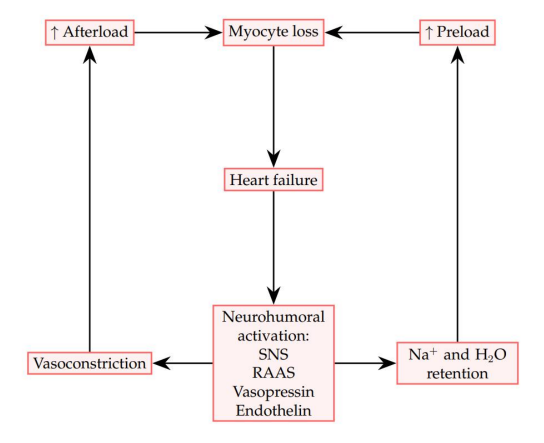


Figure 2.8: Neurohumoral activation and compensatory mechanisms in HF

- o Tachycardia
- o Complete heart block

## **▶** Diastolic dysfunction

- o Constrictive pericarditis
- o Cardiac tamponade
- o Hypertrophic / restrictive cardiomyopathies
- o LVH and fibrosis

#### **Features**

#### **Acute LVF**

- ► Sudden onset **dyspnoea** at rest
  - o Progresses rapidly to respiratory distress, orthopnoea
- **▶** Cool peripheries
- ► Rapid pulse
- ► ↑ BP (due to SNS activation)

- o May be low or normal if goes into cardiogenic shock
- ▶ ↑ JVP esp. if associated with fluid overload / RVF
- ▶ **Gallop rhythm** (tachycardia + 3 heart sounds (addition of  $S_3$ ))
- ▶ Bilateral basal crepitations if pulmonary oedema

#### **Chronic LVF**

- ► Fatigue, ↓ effort tolerance: due to ↓ CO
- ▶ Cold peripheries
- **▶** ↓ BP
- ▶ Oliguria, uraemia: due to ↓ renal perfusion
- ► Pulmonary oedema
- ► Congestive cardiac failure: RVF + LVF (RVF 2° to LVF)
  - o ↑JVP
  - o Ankle oedema (if ambulant) / sacral oedema (if bedbound)
  - Tender hepatomegaly
  - o Ascites
  - o Pleural effusion

## **Complications**

- ► Renal failure
- ▶ Hypo-K: due to
  - o Treatment with loop / thiazide diuretics
  - Impaired aldosterone metabolism in congested liver
- ► Hyper-K: due to treatment with ACEi / ARB, K-sparing diuretics (spironolactone, eplerenone)
- ► Hypo-Na:
  - o Feature of severe HF; bad prognostic sign
  - Due to
    - Diuretic therapy
    - ↑ ADH secretion
- ► Impaired liver function: due to
  - Hepatic venous congestion
  - Poor perfusion
- ► Thrmboembolism
- ► Arrhythmias: due to
  - Electrolyte abnormalities (hypo-K, hypo-Mg)
  - Underlying cardiac disease
  - Sympathetic overactivity
- ▶ Sudden death: due to V-fib

## Investigation

#### ► CXR

- Distended upper lobe pulmonary veins
- Lung fields plethoric
- o Bat-wing opacities (pulmonary oedema)
- Septal / Kerley-B lines: horizontal lines in the costophrenic angles
- Cardiomegaly (usually with chronic)

## ► Echocardiography

- o Detecting the aetiology
- Assess the ejection fraction
- o Detecting asymptomatic valvular disease

#### **▶** BNP

- o Elevated
- o Prognostic marker

## Management

#### Acute

- ▶ ALVF with pulmonary oedema is a medical emergency
- ► Continuous monitoring of ECG, BP, pulse oximetry
- ► Mnemonic: **SOPNiL** 
  - Sit (propped-up position): ↓ preload
  - High-flow  $O_2$ : correct hypoxia
  - o Ensure continuous +ve airway **pressure** (CPAP): ↓ preload
  - $\circ$  **Nitrates**:  $\downarrow$  preload, afterload
  - $\circ$  **Loop diuretic**: correct fluid overload;  $\downarrow$  preload
- ▶ If the above measures are ineffective, inotropes e.g. dobutamine.

#### Chronic

#### ► General measures

- o Diet:
  - Wt reduction
  - Avoiding high salt intake
- o Cessation of alcohol consumption
- Cessation of smoking
- Moderate exercise within limits of symptoms
- o Consider influenza and pneumococcal vaccines

## ► Drug therapy

Diuretics

- ↓ preload
- Normally, reduced preload would reduce contractility as well, but in HF, the myocardium is already beyond the maximum of the Starling curve (i.e. preload vs contractility / CO)
- o ACEi / ARB:
  - Interrupt the vicious cycle of neurohumoral activation by inhibiting RAAS (Fig. 2.8)
- Neprilysin inhibitors:
  - In resistant heart failure
  - Sacubitril is the only one available currently (in combo with valsartan)
  - $\ominus$  Neprilysin  $\rightarrow \uparrow$  ANP, BNP  $\rightarrow$  diuresis, vasodilation
- Vasodilators
  - If ACEi / ARB contraindicated (e.g. bilateral renal artery stenosis)
  - Venodilators e.g. nitrates reduce preload
  - Arteriodilators e.g. hydralazine reduce afterload
- β-blockers
  - Interrupt the vicious cycle of neurohumoral activation by inhibiting SNS (Fig. 2.8)
  - ↓ arrhythmia and sudden death risk
- o Ivabradine:
  - I<sub>f</sub> channel blocker
  - ↓ HR
- o Digoxin: rate control if coexisting A-fib
- Amiodarone: if symptomatic arrhythmia present

#### ► Non-pharmacological therapy

- o Implantable cardiac defibrillators (ICD): improve prognosis if symptomatic ventricular arrhythmias with HF
- o Resynchronisation devices (pacers): if marked conduction delay
- o Coronary revascularisation: CABG / PCI
- o Cardiac transplantation:
  - Most common indications: MI, DCM (dilated cardiomyopathy)
  - Reserved for young patients with severe disease despite optimal therapy

## 2.12 Peripheral artery disease (PAD)

## **Pathogenesis**

- ▶ Risk factors: same as CAD
- ▶ In developed countries, almost all PAD is due to atherosclerosis
- ▶ PAD in DM patients associated with severe limb ischaemia

#### **Features**

ightharpoonup  $\approx 25\%$  symptomatic

- ▶ Symptomatic PAD affects legs 8x more commonly than arms
  - o In leg, superficial femoral artery most commonly involved
  - In arm, subclavian artery most commonly involved

## ► Intermittent claudication (IC)

- Most common symptom
- o Usually in calf muscles (as superficial femoral artery is the usual site of pathology)
- In thigh or buttock if iliac arteries involved
- o Pain comes on after walking upto a specific distance, rapid relief on rest

#### ► Critical limb ischaemia

- Rest pain requiring opiate analgesia and/or ulceration or gangrene present for > 2w, with ankle BP < 50mmHg</li>
- Rest pain with ankle BP > 50mmHg = SCLI (subcritical limb ischaemia)
- Severe limb ischaemia = SCLI or CLI
- o IC usually due to single segment plaque, SLI always due to multilevel disease

#### ► Acute limb ischaemia

- Due to acute thrombotic occlusion of stenotic segment / thromboembolism
- o Features: 6P
  - Pain
  - Pallor
  - Pulselessness
  - Perishing cold
  - Paraesthesia
  - Paralysis
- o Treatment:
  - Consult vascular surgeon
  - IV heparin bolus (3000-5000 U): limit thrombus propagation
  - Distinguishing thrombosis from embolism is difficult but important as management different
    - Thrombosis: treat medically with heparin, antiplatelets, statins
    - Embolism: urgent revascularisation within 6h (otherwise extensive necrosis)
  - Irreversible ischaemia → amputation

#### ► Features of chronic limb ischaemia

- o Pulses diminished / absent
- o Bruits
- o ↓temperature
- o Buerger's sign: pallor on elevation, rubor on dependency
- Muscle wasting
- o Skin, nails: dry, thin, brittle
- Loss of hair

### Investigation

- ► ABPI (ankle-brachial pressure index)
  - O Systolic ankle BP Systolic brachial BP
  - o Normal: > 1, IC: 0.5-0.9, CLI: < 0.5
- ▶ Duplex scan
- ► CT / MRI with IV contrast
- ► Intra-arterial DSA (digital subtraction angiography): for those undergoing endovascular revascularisation
- ▶ Lipid profile
- ▶ Blood sugar

### Management

- ► Medical therapy:
  - Smoking cessation
  - **Regular exercise** (30 mins of walking, 3x a week)
  - Antiplatelet agents
    - Aspirin or clopidogrel (75 mg)
    - Newer drug: Voraxapar
      - Selective PAR-1 (proteasse-activated receptor 1) antagonist
      - Combo with aspirin or clopidogrel
  - o Statins
  - o Cilostazol:
    - peripheral vasodilator
    - improves walking distance
    - consider if nonresponsive to above
  - o Diagnose and treat DM, htn, anaemia, HF
- ▶ Surgical therapy: if medical therapy outcome not good after 6mo
  - $\circ$  Angioplasty  $\pm$  stenting
  - Endarterectomy
  - o Bypass

# 2.13 Buerger's and Raynaud's

### Buerger's

- ▶ Inflammatory disease of arteries distinct from atherosclerosis
- ► Usually in **young (20-30y) male smokers**
- ► Characteristically affects distal arteries
  - Claudication in the feet
  - $\circ$  Rest pain in the fingers / toes
  - Wrist and ankle pulses absent

- o Brachial and popliteal pulses present
- ► Treatment:
  - Smoking cessation often causes remission
    - Amputation usually needed if patient continues smoking
  - Vasodilators: **prostacyclin**
  - o Sympathectomy

### Raynaud's

- ► Young (15-30y) female in temperate climates
- **▶** Doesn't cause ulceration / infarction
- ► Usually no significant pain
- ➤ Treatment:
  - o Reassurance
  - Avoiding cold exposure
  - o Nifedipine if symptoms troublesome

### 2.14 Aortic diseases

### **Aortic aneurysm**

#### Intro

- ► Abnormal dilatation of aortic lumen
- ▶ Most common in infrarenal abdominal aorta
- ► Men: women = 3:1
- ► Typically in old (65-85y)
- ► In younger if associated with Marfan's

### **Pathogenesis**

- ► Most common cause atherosclerosis
- ▶ Predisposing factors: DM, htn, hyperlipidaemia
- ► Tends to run in families
- ► Associated with **Marfan's** (autosomal dominant connective tissue disease)
  - o Marfan's also associated with aortic dissection

#### **Features**

- ► Thoracic aortic aneurysms:
  - Acute severe chest pain
  - Features of AR
  - Compressive symptoms
    - Stridor

- Hoarseness
- o Massive bleeding (if erodes into adjacent structure e.g. oeso or bronchus)
- ► Abdominal aortic aneurysms:
  - o Pain: central abdomen, back, loin, iliac fossa, groin
  - Thromboembolism
  - Compressive symptoms
    - Duodenal compression → obstruction, vomiting
    - IVC compression  $\rightarrow$  leg oedema, DVT
  - Rupture

### Investigation

- ► **USG**: investigation of choice for dx
- ► CT scan:
  - o more info about size and extent
  - o standard pre-op investigation

#### Management

- ► Surgical repair
  - o Open repair:
    - treatment of choice
    - replace aneursymal segment with prosthetic (Dacron) graft
  - Endovascular aneurysm repair (EVAR):
- ▶ If aneurysm is due to Marfan's,  $\beta$ -blockers
  - ↓ rate of dilatation, risk of rupture
- ▶ Rupture of the aneurysm mostly kills before reaching the hospital, so try preventing that
- ▶ Asymptomatic: Risks of surgery outweigh risks of rupture if  $\leq 5.5$ cm
- ▶ Symptomatic: All cases must be operated, as pain often predates rupture

#### **Aortic dissection**

#### Intro

- ▶ Breach in arterial wall → blood enters media, splitting it into 2 layers (creating a false lumen)
- ► Men: women = 2:1
- ► Peak incidence: 6-7th decade
- ► Can occur in younger if associated with Marfan's, pregnancy, trauma

### **Pathogenesis**

- ► Risk factors
  - o Htn (in 80%)
  - o Atherosclerosis
  - Coarctation
  - o Genetic connective tissue disorders:
    - Marfan's
    - Ehlers-Danlos
  - o Pregnancy (usually 3rd trimester)

#### Classification

- ► Important for management
- ► Based on involvement of ascending aorta
  - Type A
    - involves ascending aorta
    - 2/3rds of cases
    - may also extend into the descending
  - o Type B: does not involve ascending aorta

#### **Features**

- ► Pain: sudden, severe, "tearing"
  - o Type A: anterior chest pain
  - o Type B: infrascapular back pain
- ► **Hypertension** unless major haemorrhage
- ► Signs of AR
- ► **Asymmetric pulses** (brachial, carotid, or femoral)
- ► Features due to occlusion of aortic branches
  - Coronary occlusion → MI
  - Carotid occlusion → stroke
  - $\circ$  Spinal artery occlusion  $\rightarrow$  paraplegia
  - $\circ$  Renal artery occlusion  $\rightarrow$  AKI
  - o Limb occlusion → acute limb ischaemia
  - $\circ \ \ Mesenteric\ occlusion \rightarrow gut\ is chaemia$

- ► CT / MR angiography: investigation of choice
- ► CXR
  - o Broadened upper mediastinum
  - o Distorted aortic knuckle
  - Lt sided pleural effusion

### Management

- ► Initial management:
  - Pain control
  - o Antihypertensive
- ▶ Definitive treatment:
  - o Type A: emergency surgery to replace ascending aorta
  - o Type B:
    - Treated medically (↑↑↑ surgical risk)
      - Goal: maintain MAP (mean arterial pressure) at 60-75 mmHg
      - First line therapy:  $\beta$ -blocker (labetalol preferred for additional  $\alpha$ -blockade)
      - If  $\beta$ -blockers contra  $\rightarrow$  rate-limiting CCB (verapamil, diltiazem)
      - If inadequate control Na-nitroprusside
    - Unless rupture / risk of rupture / vital organ ischaemia (gut / kidney)

#### **Aortitis**

- ► Syphilis is a rare cause
  - o Characteristic saccular aneurysm of ascending aorta

### Marfan's

#### Intro

- ► Autosomal dominant connective tissue disease
- ▶ Mutations in gene FBN1 (codes for fibrillin: an ECM protein)

#### **Features**

- ► Aortic dissection
- ► Aortic aneurysm
- ► AR / MR
- ► Skin laxity
- ▶ Joint hypermobility
- ► Long arms, legs, fingers (arachnodactyly)
- ► Scoliosis, pectus excavatum
- ► High-arched palate
- ▶ Ocular abnormalities:
  - Lens dislocation
  - Retinal detachment

- ► Genetic testing: confirmatory
- ► CXR: aortic dilatation
- ► Echo: valvular disease

### Management

- $\triangleright$   $\beta$ -blockers:
  - o ↓risk of aortic dilatation
  - o given in all Marfan's patients
- ► Avoid activities that raise cardiac output
- ▶ Surgical replacement or aortic root: if progressive dilatation

#### Coarctation of the aorta

#### Intro

- ► Mostly congenital, very rarely acquired
- ► Male : female = 2 : 1
- ► Associated with:
  - Bicuspid aortic valve (normally, only mitral valve should be bicuspid, all the others tricuspid)
  - o Berry aneurysms in cerebral circulation

### **Pathogenesis**

- ▶ Narrowing most commonly where the ductus arteriosus joins the aorta (just distal to the origin of the 3 main branches: brachiocephalic, lt common carotid, lt subclavian)
  - o Proximal to narrowing (head & neck vessels)  $\rightarrow \downarrow BP$
  - o Distal to narrowing  $\rightarrow \downarrow BP$

#### Investigation

- ► MRI: investigation of choice
- ► CXR: may show changes in the contour of the aorta

#### Management

- ▶ Untreated → death due to
  - o LVF
  - Aortic dissection
  - Haemorrhagic stroke
- ► Surgical repair:
  - $\circ$  Do in early childhood  $\rightarrow$  no htn
  - $\circ$  Surgery done later  $\rightarrow$  htn persists
- ▶ Balloon dilatation / stenting: if recurrence

2.15. HYPERTENSION 41

# 2.15 Hypertension

### Management

- ► Lifestyle changes
- **▶** Drug therapy:
  - o Guidelines: figure 2.9
  - o ACEi:
    - Use with care in impaired renal function or bilateral renal artery stenosis
    - S/E:
      - 1st dose hypotension
      - Cough
      - Rash
      - Hyperkalaemia
      - Renal dysfn
  - o CCB:
    - S/E of dihydropyridines (amlodipine/nifedipine):
      - Flushing
      - Palpitation
      - Fluid retention
    - S/E of verapamil: constipation

### 2.16 Acute rheumatic fever

#### Intro

- ► Mainly in low- and mid-income countries
- ► Mainly affects children (5-15y)

### **Pathogenesis**

- ▶ Delayed immune response to group A strep infection (sore throat / skin inf)
- $\blacktriangleright$  Anti strep-A cross-react with cardiac myosin and sarcolemmal membrane proteins  $\rightarrow$  inflammation in the endo, myo, pericardium
- ▶ Also inflame joints by same mechanism
- ► Histology:
  - o Fibrinoid degeneration in the collagen of connective tissue
  - o **Aschoff nodules:** pathognomonic
    - Only in the heart
    - Multinucleated giant cells surrounded by macrophages, T cells

#### **Features**

► Revised Jones criteria

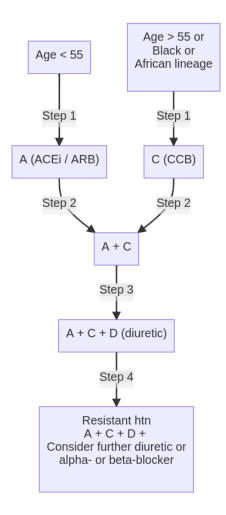


Figure 2.9: Antihypertensive therapy NICE guidelines

### o Major

- Carditis
- Polyarthritis
- Chorea
- Erythema marginatum
- Subcutaneous nodules

#### o Minor

- Fever
- Arthralgia
- ↑ ESR / CRP
- Leucocytosis
- 1° AV block
- Previous rheumatic fever

### o Supporting evidence

- Recent scarlet fever
- ↑ ASO titre

- +ve throat culture
- $ightharpoonup \ge 2$  major / (1 major +  $\ge 2$  minor) with supporting evidence
- ▶ Presumptive dx can be made without evidence of preceding strep inf if isolated chorea or pancarditis (if other causes excluded)
- **▶** Carditis
  - Manifestations:
    - Breathlessness (due to HF or pericardial effusion)
    - Palpitations
    - Chest pain (due to pan/pericarditis)
    - Pericardial friction rub (if pericarditis)
    - Soft pansystolic murmur (due to MR)
    - Soft mid-diastolic murmur (Carey Coombs murmur) due to valvulitis
    - AR (in 50% cases)

#### ► Arthritis

- o **Asymmetric, migratory, involving large joints** typically knees, ankles, elbows, wrists
- o Inflammation "migrates" from a joint within 1d 4w

#### **▶** Skin lesions

- o Erythema marginatum
  - In < 5% cases
  - Involves trunk and extremities, not face
- Subcutaneous nodules
  - In 5-7% cases
  - Small (0.5-2cm), firm, painless, over extensor surfaces
  - Typically appear  $\approx 3$  wks after the onset of other manifestations

### ► Sydenham's chorea / St Vitus dance

- o Usually appear ≥ 3mo after the episode, when all other features may have disappeared
- In  $\approx 1/3$ rd cases, more in female
- Spontaneous recovery within a few months

- ▶ Blood: CBC, ESR, CRP
- ► Throat culture
- ► ASO titre
- ► CXR: cardiomegaly, pulmonary congestion
- ► ECG: 1° AV block, features of pericarditis
- ► Echo: valvular disease

#### **Treatment**

- ▶ Bed rest
- ► Antibiotics:
  - o On dx: eliminate any residual strep infection by
    - IM benzathine benzylpenicillin (1.2M U, single dose) or
    - Oral phenoxymethylpenicillin (250mg qds 10d)
    - If penicllin allergy, erythromycin / cephalosporin
  - Long-term prophylaxis:
    - IM benzathine benzylpenicillin (1.2M U, monthly) or
    - Oral phenoxymethylpenicillin (250mg bds)
    - Continue for 5 yrs or until 21y age, whichever later
    - If residual heart disease, for 10 yrs or until 40y, whichever later

### ► Aspirin:

- o Relieves arthritis rapidly
- o Starting dose: 60 mg/kg 4hrly
- o Response within 24h helps to confirm dx
- o Continue until ESR falls
- o S/E:
  - Nausea
  - Tinnitus
  - Deafness
  - Vomiting
  - Tachypnoea
  - Acidosis
- ► Glucocorticoids:
  - o More rapid symptomatic relief than aspirin
  - o Use in carditis or severe arthritis

### 2.17 Mitral valve disease

#### Mitral stenosis

### **Pathophysiology**

- ► Almost always rheumatic
- ▶ Valve orifice narrowed by **progressive fibrosis**, **calcification of leaflets**
- ► Dilatation and hypertrophy of LA
- $ightharpoonup \uparrow LA$  pressure  $\rightarrow$  pulmonary congestion, breathlessness
- ▶ Progressive dilatation of LA commonly leads to AF
- ▶ Pulmonary htn may develop, protecting from pulmonary oedema, but leading to RVH, TR, RHF

#### **Features**

- ► Very slow progression
- ► Exertional dyspnoea: most dominant symptom
- ► **Fatigue** due to \( \psi \) cardiac output
- ► Acute pulmonary oedema → haemoptysis
- ► Thromboembolism especially with AF
- ► Exam findings:
  - $\circ$  Loud  $S_1$
  - o Tapping apex beat
  - o Low-pitched mid-diastolic murmur accentuated by exercise
  - o Basal creps if pulmonary oedema

### Investigation

- ► Doppler echo: investigation of choice
- ► ECG: AF, p-mitrale (bifid p-wave due to LA hypertrophy)
- ► CXR:
  - Straight It heart border
  - o Double rt heart border
  - o Widened carinal angle
  - May be pulmonary congestion or oedema
- ► Cardiac catheterisation if valvuloplasty considered

#### **Treatment**

- ► Medical:
  - $\circ$  **Anticoagulation:** to  $\downarrow$  thromboembolic risk
  - o **Ventricular rate control:** digoxin,  $\beta$ -blockers, rate-limiting CCB
  - o Diuretics: to relieve pulmonary congestion
- ► Valvuloplasty:
  - o Treatment of choice if
    - Significant symptoms
    - Isolated mitral stenosis with no/trivial mitral regurgitation
    - Mobile, non-calcified valve on echo
    - No thrombus in lt atrium
  - o Procedure: illustrated in fig. 2.10
  - Follow up yearly as risk of restenosis

### Mitral regurgitation

### **Pathogenesis**

► Causes:

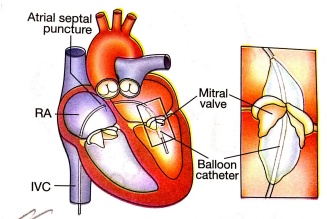


Fig. 16.82 Mitral valvuloplasty. A guidewire is introduced into the right atrium (RA) from the femoral vein and the inferior vena cava (IVC). The inter-atrial septum is punctured, providing access to the left atrium and mitral valve. A balloon catheter is then advanced over the guidewire across the mitral valve and the balloon dilated to stretch the valve and reduce the degree of stenosis.

Figure 2.10: Mitral valvuloplasty

- **Rheumatic disease** is the main cause in countries where rheumatic fever is common (causes damage to the valve cusps and chordae)
- o Mitral valve prolapse
- o Ischaemia or infarction of the papillary muscles
- o MI
- Dilated cardiomyopathy
- o Endocarditis

#### **Features**

- **▶** Breathlessness
- **▶** Fatigue
- ► Oedema, ascites (if RHF)
- ► Palpitation (if AF)
- ► Exam findings:
  - $\circ$  Soft  $S_1$
  - o Apical pansystolic murmur radiating to axilla
  - o Creps
  - o ↑JVP

- ► Echo: condition of wall and valves
- ▶ **Doppler:** quantify regurgitation
- ▶ ECG: AF due to atrial dilatation
- ► CXR:
  - o Enlarged LA, LV

- o Pulmonary venous congestion
- o Pulmonary oedema

#### Management

- ► Medical:
  - o Diuretics
  - Vasodilators if htn
  - Digoxin and anticoagulants if AF
- ► Surgical valve repair: treatment of choice for severe MR

### 2.18 Aortic valve disease

### **Aortic stenosis**

### **Pathogenesis**

- **▶** Causes:
  - o Congenital AS
  - Calcification of congenitally bicuspid aortic valve
  - o Rheumatic AS
  - Senile degenerative AS
- ► Stenosis → progressive LVH → angina (mismatch of myocardial demand and coronary arterial supply) → LVF → pulmonary oedema

#### **Features**

- ► Cardinal symptoms: angina, breathlessness, syncope
- ► Exam findings:
  - Ejection systolic murmur like "saw cutting wood"
  - o Slow-rising carotid pulse
  - o Heaving apex beat due to LV pressure overload
  - o Narrow pulse pressure
  - Pulmonary oedema
- ▶ Asymptomatic for many years; deteriorate rapidly when symptoms develop
- ▶ Die within 5y of presentation if untreated

- ► Echo: see ventricular wall, valve
- **▶ Doppler:** severity of stenosis
- ► ECG:
  - o LVH
  - "Strain" pattern: down-sloping ST segments and T-inversion in lateral leads (due to LV fibrosis)

- o LBBB
- ► CXR: LV enlarged

#### **Treatment**

- ► Asymptomatic:
  - Conservative mx
  - Regular review with Doppler (yearly; if older, 6 monthly)
- ► Symptomatic:
  - o Prompt aortic valve replacement
  - $\circ$  Delay  $\rightarrow \uparrow$  risk of sudden death
  - o Old age is not contraindication to surgery
  - o Balloon valvuloplasty: if congenital AS
  - o Anticoagulants if AF / if valve replacement with mechanical prosthesis

### Aortic regurgitation

### **Pathogenesis**

- ► Causes:
  - o Congenitally bicuspid aortic valve
  - o Rhematic disease
  - Infective endocarditis
  - o Trauma
  - Causes of aortic dilatation
    - Marfan's
    - Aneurysm
    - Aortic dissection
    - Syphilis
    - AnkSpond #### Features
- ▶ Mild to moderate: asymptomatic, palpitations
- ► Severe: breathlessness, angina
- ► Exam findings:
  - High volume/collapsing pulse
  - o ↑ pulse pressure
  - o Early diastolic murmur best heard at lt sternal edge
  - **Soft mid-diastolic "Austin-Flint" murmur**: due to regurgitant jet partly closing mitral leaflet (functional mitral stenosis)

- ▶ **Doppler:** first choice
- ► ECG: LVH, T-inversion
- ► CXR: cardiomegaly, aortic dilatation, LVF
- ► Cardiac cath

#### **Treatment**

- ► Treat underlying disease (e.g. endocarditis / syphilis)
- ► Asymptomatic: annual follow up with echo
- ▶ Valve replacement if symptomatic
- ► Vasodilators if htn

# 2.19 Tricuspid valve disease

### Tricuspid stenosis

### **Pathogenesis**

- ▶ Usually rheumatic; almost always in association with mitral and aortic valve disease
- ► TS and TR may also occur in carcinoid

#### **Features**

- ▶ ↑JVP with prominent a wave, slow y descent
- ► Mid-diastolic murmur best heard at lower rt sternal edge
  - o Higher pitch than MS murmur
  - ↑ on inspiration
- ▶ RHF → hepatomegaly, ascites, peripheral oedema

### Investigation

**▶ Doppler echo** confirms dx

#### **Treatment**

- ► Valvotomy / replacement
- ► Balloon valvuloplasty if isolated TS

# 2.20 Pulmonary valve disease

### **Pulmonary stenosis**

- ► Usually congenital
- ► May occur with carcinoid syndrome
- ► Features:
  - o Ejection systolic murmur loudest at lt uppper sternal edge radiating to lt shoulder
  - $\circ$  Wide split  $S_2$
  - o May be thrill, best felt when leaning forwards, breathes out
  - o RVH
- ► Investigation:
  - o **Doppler echo:** definitive investigation

- o ECG: RVH
- o CXR: post-stenotic pulmonary dilatation
- ► Management:
  - Mild-moderate: no treatment required
  - o Severe: balloon valvuloplasty / surgical valvotomy

### 2.21 Prosthetic valves

- ▶ 2 types:
  - Mechanical: generate prosthetic heart sounds
  - Biological: generate normal heart sounds
    - Pig
    - Allograft
- ► All mechanical valves demand long-term anticoagulation as ↑↑↑ risk of systemic thromboembolism
- ▶ Biological valves in and of themselves don't raise the risk of thrombosis, but many undergoing valve replacement will need anticoagulation anyway for risk of recurrence of AF
- ► Mechanical valves more durable than biological

### 2.22 Infective endocarditis

### **Pathophysiology**

- ► Typically at sites of pre-existing endocardial damage
- ► Common complication of IV drug use Staph endocarditis of tricuspid valve
- ▶ Sites of endocardial damage attract platelets and fibrin → vulnerable to colonisation by blood-borne organisms
- ► Avascular valve tissue and presence of fibrin and platelet aggregates help to protect proliferating microbes
- ► Vegetations (= microbes + fibrin + platelet) eventually grow large enough to cause obstruction/embolism

### Microbiology

- ▶  $\geq$  75% cases due to Staph or Strep
- ► Common cause of acute endocarditis:
  - o Staph aureus (most common)
  - o Strep pneumo
  - Strep pyo
- ► Common cause of subacute endocarditis: Viridans strep (mitis, sanguinis)
  - o oral commensals

- o may enter the bloodstream during chewing, brushing, or dental treatment
- ▶ Common cause of prosthetic valve endocarditis: coagulase -ve Staph e.g. *Staph epidermidis*
- ► Enterococcus, Strep gallolyticus (aka Strep bovis) enter the blood from the bowel or urinary tract
  - Endocarditis caused by gallolyticus should be colonoscopied, as gallolyticus associated with large gut cancer
- ► Gram -ve HACEK group cause 3-4% cases
- ▶ Brucella endocarditis: history of cattle contact
- ► Fungal endocarditis: immunocompromised / indwelling catheter

#### **Features**

- ► Modified Duke criteria
  - Major
    - Persistent +ve blood cultures >12h apart
    - $\geq 3$  +ve cultures over >1h
    - Vegetations on echo
    - New valvular regurgitation
  - Minor
    - Predisposing valvular / cardiac abnormality
    - IV drug abuse
    - Pyrexia  $> 38^{\circ}$
    - Blood culture +ve but not reaching major criteria
    - Embolic disease
    - Vasculitic disease
  - $\circ$  2 major / (1 major + 3 minor) / 5 minor  $\rightarrow$  definite endocarditis
  - $\circ$  (1 major + 1 minor) / 3 minor  $\rightarrow$  possible endocarditis
- ► Haematuria
- ▶ Heart failure
- ▶ New murmur
- ▶ Petechial rash
- ► Splenomegaly (if long-standing)
- ▶ Finger clubbing
- ► Splinter haemorrhage
- ► Osler's nodes:
  - o Rare
  - Painful, tender fingertip swellings
  - Due to vasculitis
- ► Roth's spots on ocular fundus: rare

### Investigation

**▶** Blood culture and sensitivity

#### ► Echo

- Transthoracic or transoesophageal (TOE)
- o TOE can detect vegetations as small as 1-1.5mm
- ▶ Urinalysis: proteinuria, microscopic haematuria
- ► ECG: AV block
- ► CXR: cardiomegaly, HF

### Management

- ▶ Prompt removal of any source of infection (e.g. tooth with apical abscess)
- ► Antibiotic therapy:
  - o Subacute:
    - Withhold antibiotic treatment until C/S report
    - If empirical needed, amoxicillin IV (2g 6x daily) with/without gentamicin
  - o Acute:
    - Empirical with vancomycin IV (1g bds) + gentamicin IV (1mg/kg bds)
  - o Prosthetic valve endocarditis:
    - Vancomycin IV (1g bds) + gentamicin IV (1mg/kg bds) + rifampicin oral (300-600mg bds)
  - o Duration of therapy: 2w if fully susceptible strain
- ► Cardiac surgery with **debridement of infected material** and **valve replacement**, especially in those with *Staph aureus* or fungal endocarditis

# 2.23 Congenital heart disease

#### Intro

- ▶ Maternal rubella associated with
  - o PDA
  - o PS
  - o ASD
- ► Maternal alcohol abuse → septal defects
- ► Maternal lupus → congenital CHB (complete heart block)
- ► Central cyanosis if desaturated blood enters systemic circulation without passing through the lungs (rt to lt shunt)
- ▶ Prolonged cyanosis → finger / toe clubbing
- ► Eisenmenger syndrome:
  - o Prolonged severe L2R shunt (e.g. VSD, PDA)  $\rightarrow$  progressively  $\uparrow$  RV pressure  $\rightarrow$  reversal of shunt (i.e. L2R becomes R2L shunt)

### Patent ductus arteriosus (PDA)

#### **Features**

- ▶ Dyspnoea
- ▶ Continuous "machinery" murmur with late systolic accentuation, max in the lt 2nd ICS
- ▶ Thrill
- ▶ ↑ pulse volume
- ► May eventually progress to Eigenmenger's

### Investigation

- ► Echo: inv of choice
- ► ECG: RVH
- ► CXR: enlarged pulmonary artery

#### **Treatment**

- ► Closure by cardiac catheterisation with implantable occlusive device
  - Done in infancy if significant shunt
  - o Done in late childhood if smaller shunt (closure ↓ endocarditis risk)
- ► Prostaglandin synthetase inhibitor (indomethacin / ibuprofen) in the first wk of life to medically induce closure
  - If severe PS with L2R shunt thru ductus, ductus should be kept patent until treatment to improve oxygenation

#### Coarctation of the aorta

Discussed in subsection 2.14

### **ASD**

- $\blacktriangleright$  Male:female = 1:2
- ▶ Most are ostium secundum defects involving fossa ovalis
- ightharpoonup RA to LA shunt ightharpoonup gradual enlargement of rt heart
- ► Features:
  - o Most asymptomatic for years, detected at routine exam / CXR
  - o Dyspnoea
  - Chest infections
  - o Heart failure
  - o Arrhythmias esp. AF
  - $\circ$  Volume overload of RV  $\rightarrow$ 
    - Wide fixed split of S<sub>2</sub>
    - Systolic flow murmur over pulmonary valve
- ► Investigation:

- o Echo:
  - RV dilatation, hypertrophy
  - Pulmonary artery dilatation
- TOE: precise size and location of defect
- o CXR: cardiomegaly, pulmonary plethora
- ECG: incomplete RBBB
- ➤ Treatment:
  - $\circ$  If pulmonary flow  $\geq$  1.5x systemic flow, surgical closure / cardiac catheterisation using implantable closure device
  - Else conservative treatment

#### **VSD**

- ► Most common congenital heart disease (1/500 live births)
- ▶ Most commonly defect is at the junction between the muscular and membranous parts
- ► Features:
  - o Pansystolic murmur best heart at lt sternal edge radiating all over the precordium
  - Small VSD (maladie de Roger / Roger's disease) → loud murmur
  - May progress to Eigenmenger's
- ► Investigation:
  - Doppler:
    - Helps to identify small defects that are likely to close spontaneously
    - Larger defects:
      - Monitored with serial ECG and echo: see pulmonary htn signs
      - CXR: pulmonary congestion
      - ECG: biventricular hypertrophy
- ► Treatment:
  - o Small VSD  $\rightarrow$  no treatment
  - o Cardiac failure in infancy → digoxin, diuretics
  - o Persistent failure / ECG/echo suggesting pulmonary htn  $\rightarrow$  surgical repair
  - o Eisenmenger:
    - Surgical repair contraindicated
    - Heart-lung transplant only effective treatment

# 2.24 Myocarditis

#### Causes

- ▶ Viral: most common
  - Coxsackie
  - o Influenza A, B

2.24. MYOCARDITIS 55

- ► Bacterial:
  - o Borrelia burgdorferi (Lyme disease)
  - Mycoplasma pneumoniae
- ➤ Protozoal:
  - o Trypanosoma cruzi (Chagas')
  - Toxoplasma gondii
- ► Fungal: *Aspergillus*
- ► Parasitic: *Schistosoma*
- ▶ Drugs/toxins:
  - o Alcohol, cocaine
  - Anthracyclines
  - Clozapine
  - o Lithium
- ► Autoimmune:
  - o SLE
  - o RA
  - o Systemic sclerosis
  - o Sarcoidosis

#### **Features**

- ► Self-limiting in most cases
- ▶ Death may be due to ventricular arrhythmia / rapidly progressive HF
  - Cause of sudden, unexpected death of young athletes
- ▶ Present in 4 varieties:
  - o Fulminant: follows viral prodrome, results in severe HF / cardio shock
  - Acute
  - o Chronic active
  - Chronic persistent

### Investigation

- ▶ Dx made after other common causes of cardiac dysfunction excluded
- ► Echo: LV dysfunction
- ► Cardiac MRI: myocardial inflammation
- ▶ Blood for tropo-I, tropo-T, creatine kinase

### **Treatment**

- ► Mainly supportive
- ► Treat HF and arrhythmia
- ► Avoid intense exertion (may induce fatal arrhythmia)
- ► Glucocorticoids / immunosuppressants don't help

# 2.25 Cardiomyopathy

# Dilated cardiomyopathy (DCM)

### **Pathogenesis**

- ► ↑LV mass with normal/reduced thickness
- $\triangleright \ge 25\%$  are autosomal dominant
  - Mutations affecting myocyte cytoskeleton proteins (e.g. dystrophn, lamin A and C, emerin, metavinculin)
- ► X-linked muscular dystrophies (e.g. DMD) are also associated with DCM
- ▶ Advanced HIV / Late autoimmune reaction to viral myocarditis may also cause DCM

#### **Features**

- ▶ Sporadic chest pain
- ► HF
- ► Arrhythmia
- ► Thromboembolism
- ► Sudden death

### Investigation

- ▶ D/D includes ventricular dysfunction due to CAD. Dx DCM only after excluding CAD
- ► Echo
- ► Cardiac MRI
- ▶ Genetic testing if  $\geq 1$  family members affected

#### **Treatment**

- ► Control HF
- ▶ Reduce risk of sudden arrhythmic death by  $\beta$ -blockers + ACEi/ARB

# Hypertrophic cardiomyopathy (HCM)

#### Intro

► Most common form of CM (1/1000)

#### **Pathogenesis**

- ▶ Usually autosomal dominant, high penetrance, variable expression
- ► Extensive LVH + myocardial fibrosis
- ► May be generalised or confined to the interventricular septum (asymmetric septal hypertrophy)
- ▶ Stiff Lv  $\rightarrow \downarrow$ impaired diastolic filling  $\rightarrow$  HF
- ► Septal hypertrophy → dynamic outflow tract obstruction (hypertrophic obstructive cardiomyopathy, HOCM)

#### **Features**

- ▶ Very similar to AS, except jerky pulse in HCM
- ► Angina on effort
- ▶ Breathlessness on effort
- ► Arrhythmia → sudden death (typically after vigorous exertion)
- ▶ Most common cause of sudden death in young athletes
- ► Heaving apex beat
- ▶ Double impulse at apex (palpable S<sub>4</sub> due to LA hypertrophy)
- ► MDM at base
- ► PSM at apex (due to MR)

### Investigation

- ► Echo: inv of choice
- ► ECG: LVH
- ▶ Genetic testing

#### **Treatment**

- $\triangleright$   $\beta$ -blockers, rate-limiting CCB, disopyramide: relieve symptoms, prevent syncope
- ► Amiodarone: control arrhythmia
- ► Relief of outflow obstruction by
  - o Partial myectomy
  - o Catheter ablation of basal septum
  - o Avoid digoxin, vasodilators (may ↑ outflow obstruction)

### 2.26 Cardiac tumours

- ▶ 1° cardiac tumours rare
- ► Most (75%) 1° are benign
  - o Myxoma
  - o Fibroma
  - o Lipoma
  - o Fibroelastoma
  - Haemangioma
- ► Atrial myxoma: most common in LA, treat by surgical excision

# 2.27 Pericardial diseases

► Normal pericardial fluid: 50 mL

### Acute pericarditis

### **Pathogenesis**

► Causes:

- o Infection: viral, bacterial, tubercular
- o Inflammatory: RA, RF, SLE
- o Post-MI
- o Malignancy
- ► All forms may produce pericardial effusion
  - Haemorrhagic effusion → ca breast / bronchus / lymphoma

#### **Features**

- ► Retrosternal chest pain
  - Radiates to shoulders and neck
  - o Aggravated by deep breathing, movement, change of posture, exercise, swallowing
- ▶ Low-grade fever
- ► Pericardial friction rub:
  - o Diagnostic of pericarditis
  - o High-pitched, scratching / crunching noise
  - Typically during systole

### Investigation

- ▶ Dx by features + ECG
- ▶ ECG: ST-elevation with upward concavity in leads over the affected area

#### **Treatment**

- ► Pain relief:
  - o Aspirin (600mg, 6x daily)
  - o Indomethacin (50mg tds)
- ▶ Viral pericarditis: recover within a few days/wks
- ► Purulent pericarditis: antibiotic therapy

#### Pericardial effusion

#### **Features**

- ▶ Often accompanies pericarditis
- ▶ Quieter heart sounds
- ► Friction rub diminished in intensity
- ▶ Large / rapidly developing effusion → cardiac tamponade (acute HF due to compression)
  - o Dyspnoea
  - ↑↑↑ JVP
  - o Kussmaul's sign: paradoxical ↑ JVP during inspiration
  - o Hypotension
  - o Tachycardia

- Pulsus paradoxus (abnormally large decrease in stroke volume, systolic blood pressure and pulse wave amplitude during inspiration. The normal fall in pressure is less than 10 mmHg.)
- Oliguria

### Investigation

- ► Echo: inv of choice
- ► CXR:
  - o Enlarged cardiac silhouette
  - Large effusion → globular heart shadow

#### **Treatment**

▶ Large effusion causing tamponade (medical emergency) → pericardiocentesis under echo guidance

### **Tuberculous pericarditis**

#### **Treatment**

► Anti-TB therapy + 3mo prednisolone course

### Chronic constrictive pericarditis

- ▶ Progressive thickening, fibrosis, calcification of pericardium
- ▶ Heart surrounded by solid shell → impaired filling

#### **Features**

- ► S/S of systemic venous congestion
- ► Fatigue
- ► Rapid, low-volume pulse
- ▶ †JVP
- ► Kussmaul's sign
- ▶ Venous congestion → hepatomegaly, ascites, peripheral oedema
- ► Pulsus paradoxus

### Investigation

- ► CXR: pericardial calcification
- ► Echo
- ► CT scan

#### **Treatment**

ightharpoonup Diastolic HF ightharpoonup loop diuretics, aldosterone antagonists

# Chapter 3

# Dermatology

# 3.1 Anatomy and physiology

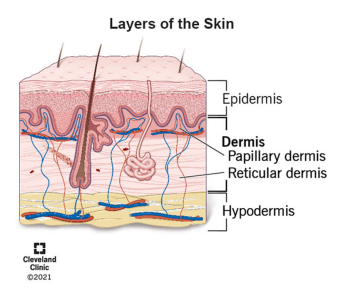


Figure 3.1: Layers of the skin

### **▶** Layers of skin:

- o **Epidermis**: further layered into (from out→in)
  - corneum
  - lucidum: only in thick skin i.e. palm and sole
  - granulosum
  - spinosum
  - basale
- o Dermis:
  - Papillary dermis: more superficial part, containing
    - loose areolar tissue
    - subpapillary vascular plexus

- Reticular dermis: deeper part, containing
  - collagen
  - deep vascular plexus
  - nerves
  - pilosebaceous units (hair follicle + sebaceous gland)
  - sweat glands
  - some keratinocytes around the hair follicles and sweat glands (important for wound healing - these are the reasons behind healing without scarring in upto partial-thickness skin wounds)
- o Subcutis / subcutaneous tissue: (technically not part of the skin) adipose tissue

### **Epidermal appendages**

- ► Hair follicles:
  - o phases of growth
    - anagen:
      - active growth
      - lasts years in scalp hairs
    - catagen:
      - transitional
      - lasts days (in scalp)
    - telogen:
      - resting
      - lasts months (in scalp)
- ► Sebaceous glands
  - o usually associated with a hair follicle (called a pilosebaceous unit)
  - o  $| androgens | \rightarrow \uparrow | sebum$  o  $| oestrogen | \rightarrow \downarrow | sebum$
- ► Sweat glands
  - o innerved by *sympathetic cholinergic* fibres

# 3.2 Principles of management of skin disease

### **Topical treatments**

- ► Ointments vs Creams
  - o Ointments preferred to creams for dry skin (e.g. chronic eczema) as
    - more hydrating
      - 80% oil + 20% water in ointments (vs 50-50 for creams)  $\rightarrow$  prevent water loss from skin by oil layer
    - less preservatives  $\rightarrow$  less risk of allergy

- **▶** Emollients
  - o Moisturise, lubricate, protect skin
  - Vehicles without active drug
- ► Gluocorticoids

### **Phototherapy**

- ► UVB
- ► Psoralen UVA
  - o Psoralen:
    - natural photosensitiser from plant source
    - cross-link DNA strands on excitation with UVA
  - $\circ$  Cumulative exposure to PUVA  $\rightarrow \uparrow$  risk of SCC, so reserved for UVB resistance
- ▶ Uses
  - o Psoriasis
  - o Atopic eczema
  - o Vitiligo
  - o Chronic urticaria

### **Systemics**

- ► Antihistamines
- **▶** Retinoids
  - o Anti-inflammatory
  - o Promote differentiation of skin cells
  - o Teratogenic
    - must be prescribed with robust contraception
    - females must have negative pregnancy test before, during, and after therapy

### **▶** Immunosuppressants

- o Glucocorticoids e.g. prednisolone
- Methotrexate
- Azathioprine

# **Biologics**

- ▶ Biological inhibitors of proinflammatory cytokines
- ▶ TNF- $\alpha$  inhibitors
  - o Infliximab
  - o Etanercept
- **▶** Interleukin inhibitors
  - o Ustekinumab: IL-12, 23

3.3. SKIN CANCERS 63

- Guselkumab: IL-23Secukinumab: IL-17
- ► *Rituximab*:
  - o Binds to CD20  $\rightarrow$  cause ADCC of B cells
  - o As terminally differentiated plasma cells don't have CD20 they're safe
  - Use: pemphigus vulgaris

### Non-surgical therapy

- ► Cryo
  - o Liquid N<sub>2</sub>
  - o Causes cell membrane destruction → death
- ► Laser
- ► PDT / photodynamic therapy

### 3.3 Skin cancers

#### Classification

- ▶ Non-melanoma skin cancer (NMSC): most common
  - o SCC
  - o BCC
- ▶ Melanoma
  - Less common
  - o More metastatic risk  $\rightarrow$  cause of most skin cancer deaths

# 3.4 Fungal infections

### **Types**

- ► Superficial
  - o Dermatophytes: aka ringworm / tineasis
    - Trichophyton
    - Epidermophyton
    - Microsporum
  - o Yeast
- ► Deep: less common
  - o Chromomycosis
  - Sporotrichosis

### 3.5 Scabies

### Agent

Caused by the mite Sarcoptis scabies hominis

### **Diagnosis**

- ► Identify the skin burrow
- ▶ Visualize the mite by dermatoscope / extracting with a needle

#### **Treatment**

- ► Affected + all asymptomatic family members / physical contacts
- ► Topical permethrin / malathion
  - o 2 applications
  - o 1 wk apart
  - Whole body, except head
- ► Oral Ivermectin:
  - o Single dose
  - o For poor adherence, immunosuppresion or heavy infestation

### 3.6 Acne

► Chronic inflammation of pilosebaceous units

## **Pathogenesis**

Key components are:

- ► ↑ Sebum production
- ► Colonisation of pilosebaceous ducts by *Propionibacterium acnes*
- ► Occlusion of pilosebaceous ducts

#### **Features**

- ► Hallmark: comedone
- ► Greasiness of skin

### Management

- ► Mild disease
  - o Topical Benzoyl peroxide
  - o Topical Retinoids
  - Topical antibiotics

3.7. ECZEMAS 65

- Erythromycin
- Clindamycin
- ► Moderate disease: topical *plus* 
  - Systemic tetracycline
  - Oestrogen containing OCP
  - o Isotretinoin: if inadequate response to topical+systemic therapy for 6 months

#### **▶** Severe disease

- o Isotretinoin 0.5-1 mg/kg for 4 months:
  - Reduce sebum secretion and follicle colonisation
  - Teratogen
  - Pregnancy must be avoided during treatment and within 2 mo of drug cessation
- Systemic glucocorticoid (with isotretinoin)
- If unable to use isotretinoin
  - UVB phototherapy
  - PDT

### 3.7 Eczemas

▶ Seborrhoeic dermatitis is associated with *Malassezia* yeasts

#### **Features**

Most types have the following clinical features:

#### Acute

- ► Ill-defined erythema, oedema
- ▶ Papules, vesicles, bullae
- **▶** Exudation
- ▶ Scaling

#### Chronic

- ► Above features
- ► Lichenification
  - Skin thickening with pronounced skin markings, 2° to chronic scratching
  - o Fissures
  - Dyspigmentation

### Management



Figure 3.2: Management of eczema

# 3.8 Psoriasis

- ► Chronic inflammatory hyperproliferative skin disease
- **▶** Characteristics
  - o Well-defined erythematous scaly plaques
  - o Affecting extensor surfaces, scalp, nails

### Histological features

- ► Keratinocyte hyperproliferation + abnormal differentiation → nucleated stratum corneum cells (transit time from basale to corneum reduced to 5 from 28 → keratinocytes reach the surface while immature)
- ▶ Inflammation with Th-1 and Th-17 infiltration
- ► Tortuosity of dermal capillaries and release of VEGF

### **Exacerbating factors**

► Sunlight

3.8. PSORIASIS 67

- **▶** Trauma
- **▶** Infection
  - o  $\beta$ -haemolytic strep  $\uparrow$  guttate psoriasis
  - o HIV may initally present with severe psoriasis

### **▶** Drugs

- o Antimalarials
- o  $\beta$ -blockers
- o Lithium
- o NSAIDs
- ► Stress and anxiety

### Clinical types

- ▶ Plaque psoriasis:
  - o most common
  - o well-demarcated erythematous plaques
  - o silver-white scales in untreated
    - bleed on scraping (due to dilated vessels underneath) → **Auspitz sign**
  - o Sites
    - extensor surfaces
      - elbows
      - knees
      - lower back
    - scalp
    - nails
- ► **Guttate** psoriasis:
  - o follows *Strep* throat
  - o common in children/adolescent
  - o UVB highly effective
  - o may herald the onset of plaque psoriasis in adulthood
- **▶ Erythrodermic** sporiasis: generalised → medical emergency
- ► Pustular psoriasis

### Management

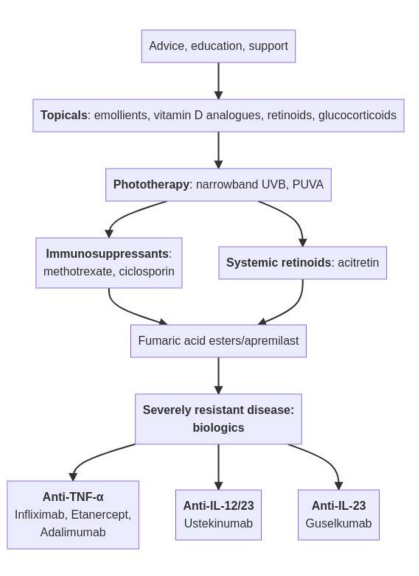


Figure 3.3: Management of psoriasis

# **Complications**

- ► Psoriatic arthropathy
- ► Exfoliative dermatitis
- ► Secondary infection
- ► Hyperuricaemia and gout

# 3.9 Hypopigmentation

### **Causes**

- ▶ Vitiligo
- ► Albinism
- ▶ Pityriasis alba
- ► Pityriasis versicolor

### Vitiligo

- ► Acquired
- ► Cell-mediated autoimmune destruction of melanocytes
- ► Loss of melanocytes → hypopigmented patches

#### Albinism

- ► Autosomal recessive
- ▶ Reduced melanin production by normal number of melanocytes
- ▶ ↑↑ risk of sunburn, skin cancer

# 3.10 Hyperpigmentation

#### **Causes**

- **▶** Endocrine
  - o Melasma/chloasma:
    - in pregnancy / some OCP users
    - discrete patches of facial pigmentation
  - o Addison's disease
  - Cushing's syndrome
  - o Nelson's syndrome
    - hyper-ACTH 2° to bilateral adrenalectomy for Cushing's
    - due to loss of -ve feedback from plasma cortisol
  - o CKD

#### **▶** Drugs

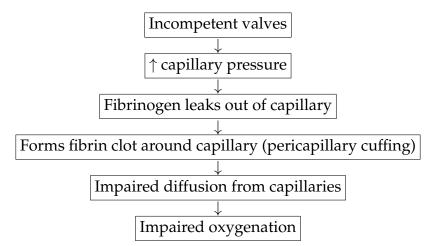
- o Amiodarone
- o Anti-cancers:
  - Bleomycin: Hodgkin's
  - Busulfan: CML
- Choroquine
- o Psoralens

# 3.11 Stuff not large enough to devote a section to

### SPF (sun protection factor)

► UV dose for producing erythema with sunscreen UV dose for producing erythema without sunscreen

### Mechanism of venous ulceration



# Chapter 4

# Nephrology

# 4.1 Anatomy and Physiology

### Nephron

- ► Glomerulus (fig. 4.1)
  - o Glomerular capillaries enclosed by Bowman's membrane
  - Space between adjacent capillaries occupied by mesangial cells
  - o Glomerular capillaries:
    - Endothelial cells with
      - fenestrations ( $\approx 70$ nm)
      - charged glycocalyx (mostly -ve)
    - GBM (glomerular basement membrane) made of type IV collagen etc
  - o **Podocytes / visceral epithelial layer:** interdigitating foot processes cover up the fenestrations, making the filtration barrier tighter
  - $\circ$  **Mesangial cells:** contract and relax to control capillary diameter  $\rightarrow$  GFR
- ► Renal tubules
  - o Lining different according to location and function
  - o Interstitial fibroblasts produce erythropoietin

# Juxtaglomerular (JG) apparatus

- ▶ Macula densa around the distal tubules: Sense Na concentration in distal tubule filtrate, controlling the tubuloglomerular feedback mechanism
  - o If low GFR, release PGE2  $\rightarrow$  dilate afferent arteriole  $\rightarrow \uparrow$ GFR
  - $\circ$  If high GFR, make adenosine from ATP  $\to$  afferent arteriolar constriction  $\to \downarrow$  GFR
    - Aside: this is the mechanism behind the mild diuresis produced by caffeine. Adenosine carries the signal for tiredness in the brain, and caffeine blocks adenosine receptors. In the brain, the blocking means you feel less tired. In the afferent arterioles, blocking means you can't reduce your GFR.
  - o If high GFR, stimulate JG cells to secrete renin

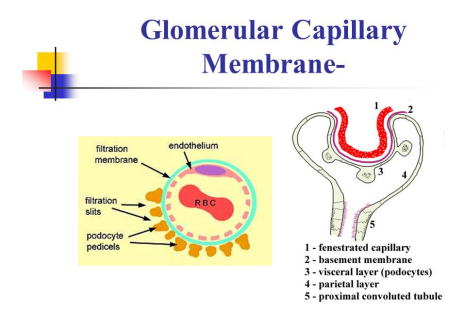


Figure 4.1: Microstructure of the glomerulus

# 4.2 Presenting problems in urinary disease

➤ Oliguria: < 400mL/day</li>➤ Anuria: < 100mL/day</li>

► Haematuria:

- o Healthy indiviuals may have upto 12,500 RBCs/mL
- Macroscopic or microscopic-dipstick+ haematuria indicates significant pathology
- o Types:
  - Initial:
    - blood during the start of micturition
    - cause: penile urethral pathology
  - Terminal:
    - blood at the end of micturition
    - cause: bladder / prostatic urethral pathology
  - Intermittent:
    - cause: IgA nephropathy, Alport's, PKD, renal tumour

#### ► Proteinuria:

- Very small amounts of high molecular weight and moderate amounts of low molecular weight proteins pass through the healthy GBM (glomerular basement membrane)
- o Whatever passes is almost completely reabsorbed by receptors on tubular cells
- So in healthy individuals, < 150mg/day protein excreted through urine
- o Transient proteinuria

- Causes
  - Vigorous exercise
  - Fever
  - Heart failure
  - UTI
- Retest after trigger has resolved to verify if persistent proteinuria
- Proteinuria best tested on early morning sample, as some have orthostatic proteinuria (benign, < 1g/day, associated with upright posture)
- $\circ$  Dipstick positive when > 0.5 g/day
  - If persistent dipstick proteinuria, 24h urinary protein must be quantified. > 1g/day → likely glomerular disease
  - PCR (Protein:Creatinine ratio) in spot sample might give better estimates than 24h protein as 24h urine collection is often inaccurate
  - Renal biopsy to confirm dx in significant proteinuria

# 4.3 Glomerular diseases ("Glomerulonephritides")

#### Introduction

- ► Though strictly means inflammation of the glomeruli, used to describe all glomerular disease (even if non-inflammatory e.g. minimal change disease)
- ► Lie on a *spectrum*, from *nephrosis* (podocyte pathology → proteinuria) to *nephritis* (inflammation + GBM damage → haematuria)

# **Types**

#### Nephrotic presentation

- ► Minimal change disease
  - Normal except on electron microscopy
  - Electron microscopy shows fusion of podocyte foot processes
- ► Focal segmental glomerulosclerosis (FSGS)
- ► Membranous nephropathy

# Mild glomerulonephritic presentation

- ► IgA nephropathy
- ► Mesangiocapillary glomerulonephritis

# Rapidly progressive glomerulonephritic presentation

- ► Focal necrotising glomerulonephritis
- ▶ Diffuse proliferative glomerulonephritis
- ► Anti-GBM disease (aka Goodpasture's syndrome)

# 4.4 Nephrotic syndrome <sup>1</sup>

#### **Features**

- ► Massive **proteinuria** (> 3.5 g/day (medicine) or  $1 \text{ g/m}^2$ /day (paediatrics))
  - o **Hypoalbuminaemia** (< 3 g/dL)
  - Generalised oedema (pitting)
  - o "Effusions": Ascites, pleural effusion, pericardial effusion
- ► Features of reduced circulatory volume
  - Scanty urination (colour normal)
  - o Pulse: weak
  - o BP: low
  - Capillary refill: prolonged (> 3s)
- ▶ Hyperlipidaemia & lipiduria
  - o due to ↑ lipoprotein production by liver
- **▶** Recurrent infections
  - due to loss of immunoglobulins with urine
- ► Features of **complications** 
  - Shiny abdominal wall, rigidity and tenderness, absent bowel sounds → peritonitis, likely by *Strep pneumo*
  - o Loin pain  $\pm$  palpable kidney + haematuria  $\rightarrow$  **renal vein thrombosis**
  - o Alterations of consciousness, hemiplegia → **stroke**

# Histopathology

- ► Histological types include
  - o Minimal change disease:
    - no pathology visible with light microscope, visible pathology only under electron microscope.
    - most common type in children (2-8y)
  - o Membranous nephropathy: most common type in adults

# Aetiopathogenesis

### Minimal change disease

# **▶** Effacement of podocyte foot processes

<sup>&</sup>lt;sup>1</sup>Most parts of this section, unless specifically mentioned to be for membranous nephropathy, refers to minimal change disease i.e. (most cases of) paediatric nephrotic syndrome

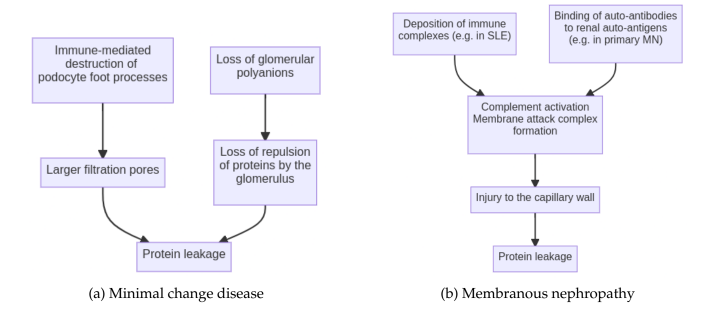


Figure 4.2: Pathogenesis of nephrotic syndrome

#### Membranous nephropathy

- ▶ Autoantibodies to podocyte surface antigens (e.g. M-type phospholipase  $A_2$  receptor 1)→ MAC-mediated glomerular capillary injury
  - Antibodies to phospholipase A<sub>2</sub> (PLA2Rab), thus, can be used for diagnosis without biopsy (but may be absent in early disease so biopsy might still be needed)

### ► Aetiology:

- o Primary / Idiopathic: HLA-DQA1
- Secondary:
  - Drugs e.g. NSAIDs, penicillamine (treatment for Wilson's)
  - Heavy metal poisoning
  - Hep-B
  - Malignancy
  - SLE

#### ► Fates:

- o 1/3 spontaneous remission
- o 1/3 remain nephrotic
- o 1/3 progressive CKD

# Investigation

- ▶ Urine RME
  - o Albuminuria
  - o Granular & hyaline casts

- o Pus cells if associated with UTI
- ► Diagnostic
  - 24h total protein: > 1g/m²/day
  - PCR (Protein-Creatinine ratio) > 2
  - Serum albumin < 25 g/L
  - Serum cholesterol > 220 mg/dL
- ► Renal biopsy: if nonresponsive to corticosteroid therapy

#### **Treatment**

### Minimal Change Nephropathy

- ► Specific:<sup>2</sup>
  - High dose glucocorticoid for 6wks
  - o Prednisolone 1 mg/kg/day
  - KDIGO guidelines: (4wks daily + 4wks alternate day) or (6wks daily + 6wks alternate day)
  - Incomplete response to steroid: Cyclophosphamide (1.5-2 mg/kg/day) or Ciclosporin (3-5 mg/kg/day) for 8-12wks with prednisolone 7.5-15 mg/day (i.e. low dose)

### **▶** Supportive:

- o Diet:
  - normal with adequate protein
  - salt restriction
- o Oedema: if severe,
  - restrict salt and fluid intake
  - loop diuretics
- Hypovolaemia: infuse albumin
- o Infections: treat & prevent (pneumococcus, meningococcus vaccine)
- Hypercoagulability:
  - due to loss of coagulation inhibitors (antithrombin III, protein C, protein S) + ↑ liver production of procoagulants
  - consider LMW heparin / warfarin if severe NS

### Membranous Nephropathy

- ► Specific:<sup>3</sup>
  - o High-dose glucocorticoids + cyclophosphamide (Ponticelli regimen)
    - Reserved for severe cases for risk of toxicity of this regimen

<sup>&</sup>lt;sup>2</sup>https://kdigo.org/wp-content/uploads/2017/02/KDIGO-GD-Guideline-Key-Takeaways-for-Clinicians-Nephrotic-Syndrome-in-Children.pdf

<sup>&</sup>lt;sup>3</sup>https://kdigo.org/wp-content/uploads/2017/02/KDIGO-GD-Guideline-Key-Takeaways-for-Clinicians-Membranous-Nephropathy.pdf

- o If secondary, treat the underlying cause
- ► **Supportive:** same as above

### **Complications**

- ► Hypercoagulability → renal vein thrombosis, pulmonary embolism
  - o Dx of RVT: Doppler, CT, or MRI
  - o Rx: Heparin 6 days (5-7d), Warfarin 6 months (3-6m)
- ▶ Infections: due to loss of immunoglobulins
  - o **Pneumococcal infection** → peritonitis, septicaemia
  - Cellulitis
  - Strep infection
- ► Hyperlipidaemia → atherosclerosis
- **▶** Pleural effusion
- ► Pericardial effusion

# 4.5 Acute post-streptococcal glomerulonephritis

### **Pathogenesis**

- ▶ Occurs following sore throat or skin infection by **group-A**  $\beta$ **-haemolytic streptococcus** (nephritogenic strains: 12, 49)
- ► Antibodies against streptococcal M protein cross-react with glomerular antigens because of molecular mimicry
- ► Complement activation, inflammatory cell infiltration. Net effects:
  - o Oliguria
  - o Haematuria
  - o Hypervolaemia, hypertension, hyperkalaemia
  - o Oedema
  - $\circ \downarrow$ Renal function  $\Longrightarrow$ 
    - Azotaemia
    - Acidosis
- ► **Complications** (due mostly to acute hypertension)
  - o Acute LVF
  - o Hypertensive encephalopathy
  - Acute kidney injury

#### **Features**

- ▶ Age: 5-12y; history of strep throat/skin infection a few wks prior
- ► Scanty, smoky urine
- **▶** Puffy face

- **▶** Hypertension
- ► Features of *complications* 
  - o ALVF:
    - cough
    - respiratory distress
    - orthopnoea
    - gallop rhythm
    - bilateral basal crepitations
  - Hypertensive encephalopathy:
    - headache
    - blurred vision
    - convulsion
    - delirium
    - papilloedema
  - o AKI: anuria

### Investigation

- **▶** Evidence of nephritis
  - o Urine RME:
    - RBC, RBC casts
    - Mild proteinuria
    - Leucocytes
  - o Serum C3: ↓
  - o Serum C4: normal
- **▶** Evidence of prior strep infection
  - o ASO titre: ↑
  - o Anti-DNAse B:↑
- **▶** Evidence of complications
  - o Serum electrolytes: may show hyperkalaemia and acidosis
  - o Serum creatinine
  - o X-ray chest

#### **Treatment**

- ► Rest
- ▶ **Diet**: restrict fluid, salt, protein.
- ▶ **Diuretics**: furosemide
- ► **Antibiotics**: phenoxymethyl penicillin PO to prevent spread of remaining strep within the body
- **▶** Antihypertensive

# 4.6 Alport's syndrome

- $\blacktriangleright$  Mutation / deletion of COL4A5 on chr-X  $\rightarrow$  **defect of collagen type IV**
- ► X-linked recessive
- lacktriangle Deposition of abnormal collagen  $\rightarrow$  progressive degeneration of GBM
  - o Haematuria starts in early infancy
  - o ESRD by late teens / twenties
  - o Female carriers usually have haematuria, rarely significant renal disease
- ▶ Other basement membranes with collagen IV are also involved
  - o cochlear BM → **SNHL** (sensorineural hearing loss; especially in high-frequencies)
  - o ocular abnormalities:
    - lenticonus
    - keratoconus
    - cataracts
    - corneal erosions
- ► ACEi may slow (but not prevent) loss of renal function
- ► Might require RRT (renal replacement therapy)

# 4.7 Adult Polycystic Kidney Disease

#### Introduction

- ▶ Better known as Autosomal Dominant PKD (ADPKD)
  - There's a much rarer (1:20,000) autosomal recessive PKD (ARPKD)
- ► Prevalence 1:1000
- ► Autosomal dominant
- ▶ Small cysts lined by tubular epithelium develop from childhood, enlarge slowly
- ▶ Surrounding normal kidney tissue compressed and progressively damaged
- ▶ Mutations: PKD1 (in 85%), PKD2 (15%) (code for polycystin 1 and 2, respectively)

#### **Features**

- ► Asymptomatic initially
- ► **Hypertension** from around 20 yrs of age
- ► Either (or both) kidney may be palpable, nodular
- ► Vague **loin discomfort** due to **enlarging mass**
- ► Acute loin pain due to haemorrhage into a cyst
- ► **Haematuria** with little / no proteinuria
- ▶ About 30% have **hepatic cysts** (mostly with no liver function impairment)
- **Berry aneurysms** of cerebral vessels in  $\approx 5\%$  (may lead to SAH)
- ► **Renal failure** → features of CKD
- ► Mitral / aortic regurgitation (frequent, rarely severe)

# Investigation

- ▶ Dx is based on family history, clinical features, and USG
- ► Criteria for dx in patients with +ve family history
  - o 15-39y:  $\geq$  3 unilateral or bilateral cysts
  - o 40-59y:  $\geq$  2 cysts in each kidney (total:  $\geq$  4)
  - $\circ \ge 60$ y:  $\ge 4$  cysts in each kidney (total: ge 8)

### Management

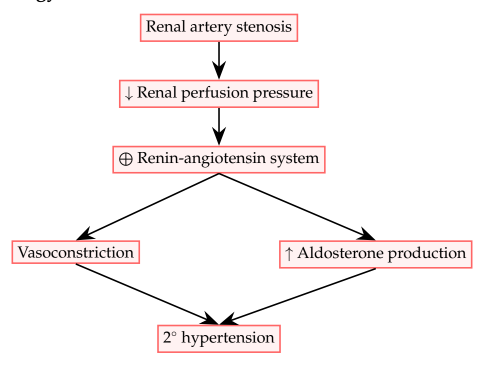
- ▶ **BP control:** 1st choice ACEi / ARBs
- ► Tolvaptan: Vasopressin V2 receptor antagonist
  - Reduce cyst growth
  - Slow rate of GFR decline
- ▶ Octreotide: Long acting somatostatin (aka growth hormone inhibiting hormone) analog
  - o Reduces rate of growth of liver and renal cysts
- ► Large cyst: US-guided aspiration / laproscopic cystectomy
- ▶ RRT (renal replacement therapy): dialysis / transplantation

# 4.8 Renal artery stenosis

#### Introduction

- ► Most common cause of 2° hypertension
- ► Most cases are due to **atherosclerosis**
- ▶ Younger ( $\leq$  40) cases mostly non-atheromatous, due to *fibromuscular dysplasia*

### **Pathophysiology**



#### **Features**

- ▶ Hypertension: may be severe and difficult to control
- ► Acute pulmonary oedema: associated with severe hypertension
- ► Progressive renal failure
- ▶ Worsened renal function with ACEi or ARB administration (≥ 30% increase in serum creatinine raises possibility of renal artery stenosis)
- ▶ Peripheral vascular disease: associated, but not caused by the stenosis
  - In older patients with atherosclerotic renal artery stenosis (due to generalised atherosclerosis)
  - o Especially in legs

# Investigation

- ► CT / MR angiogram of renal vasculature: confirmatory
- ► Serum creatinine: to see if worsening renal function
- ► Plasma renin: may be elevated
- ▶ Serum electrolytes: may be hypokalaemia due to hyperaldosteronism

► USG: asymmetrically sized kidneys

#### **Treatment**

#### Medical

- ► Antihypertensives: not ACEi or ARB
- ► Lipid-lowering agents: Statins
- ► Anti-platelets: Low-dose aspirin

#### Surgical

- Angioplasty
- ► Indications:
  - $\circ \le 40y$  age
  - o BP not controlled by antihypertensive / history of malignant htn
  - History of flash pulmonary oedema
  - Deteriorating renal function

# 4.9 Thrombotic microangiopathies (HUS, TTP)

#### Introduction

- ▶ Thrombotic occlusion of arterioles and capillaries especially in the kidneys or brain
- ► Common feature of these is microangiopathic haemolytic anaemia (MAHA)
  - Haemolysis due to passage of RBCs through abnormal vessels
  - o Schistocytes (fragmented RBC) may be seen on PBF
  - $\circ \ \ Reticulocytosis$
  - $\circ \uparrow$  unconjugated bilirubin, serum LDH
  - ↓ serum haptoglobin
    - Lysed RBCs release free Hb into the bloodstream. This can enter into tissues and trigger the inflammatory cascade, leading to tissue damage. To protect from this, the body comes equipped with a plasma protein called *haptoglobin*, which binds to free Hb. When there's intravascular haemolysis, free Hb is released, and haptoglobin is used up, leading to its decreased serum levels.

# Haemolytic uraemic syndrome

- ▶ Thrombotic microangiopathy predominantly involving renal microcirculation
- ► Cause: shiga-like toxin producing bacteria
  - o Enterohaemorrhagic Escherichia coli
  - o Shigella dysenteriae
- ▶ Most common cause of AKI in children
- ▶ If no bloody diarrhoea, atypical causes of HUS (e.g. complement system abnormalities familial HUS) should be considered

### Thrombotic thrombocytopoenic purpura

- ▶ Thrombotic microangiopathy predominantly involving cerebral microcirculation
- ► MAHA + thrombocytopoenia
- ► Autoimmune: antibodies to ADAMTS-13
  - o aka vWF cleaving protease
    - functions of vWF: haemostasis (thrombus + clot)
      - bind to and stabilize Factor VIII (unbound half-life: 1h, bound to vWF: 12h)
      - help platelet aggregation
  - o destruction of ADAMTS-13  $\rightarrow \uparrow vWF \rightarrow thrombosis$

# 4.10 Acute kidney injury (AKI)

#### Definition

- ► Sudden, (usually) reversible loss of renal function, developing over days or weeks (< 3 months), evidenced by rising serum creatinine and / or falling GFR.
- ► KDIGO criteria<sup>4</sup>: presence of any of the following
  - o  $\Delta SCr \ge +0.3 \, mg/dL \, over \, 48h$
  - o  $\Delta SCr \ge +0.5x$  baseline (or  $SCr \ge 1.5x$  baseline) over 7d
  - *Urine volume*  $\leq 0.5 \text{ mL/kg/h for} \geq 6h$

#### **Causes**

#### Pre-renal

- ▶ Cardiac failure
- ▶ Blood loss
- **▶** Dehydration
- **▶** Burns
- Sepsis

#### Renal

- ► *AGN* (acute glomerulonephritis)
- ► *AIN* (acute interstitial nephritis)
  - Drugs (more in Features)
  - Toxins
  - o Inflammatory disease
  - Infection
- ► *ATN* (acute tubular necrosis)
  - o Drugs
  - o Toxins
  - Prolonged hypotension

<sup>&</sup>lt;sup>4</sup>https://kdigo.org/wp-content/uploads/2016/10/KDIGO-2012-AKI-Guideline-English.pdf

#### Post-renal

- ▶ Bilateral renal stone
- ▶ BEP (benign enlargement of prostate)
- ► Bladder / prostate / cervical cancer
- ▶ Urethral stricture
- ▶ Meatal stenosis
- ► Phimosis

#### **Features**

- ▶ Pre-renal: features of shock
- ► Renal:
  - o AGN: haematuria, mild proteinuria, oedema, htn
  - o AIN: worsened renal function following drugs (NSAIDs, PPIs, penicillins)
    - NSAIDs  $\rightarrow \downarrow$  prostaglandins  $\rightarrow \downarrow$  afferent arteriolar vasodilation  $\rightarrow \downarrow$  GFR
  - o ATN: dense granular ("muddy brown") casts
- ▶ Post-renal: bladder distension, hydronephrosis on USG

### Management

- ► Assess fluid status
  - o If hypovolaemic, fluid resuscitation and inotropes as needed
  - Once euvolaemic, fluid intake = urine output + 500mL (for covering insensible loss)
  - o If hypervolaemic, diuretics
- ► Manage hyperkalaemia
  - $\circ$  If K<sup>+</sup> > 6.5 mmol/L
    - Calcium resonium to stabilize myocardium
    - Glucose + Insulin to reduce K
- ► Manage acidosis
  - o NaHCO₃ if pH < 7
- **▶** Treat the cause
- ► Discontinue nephrotoxic drugs
- ► Reduce doses of other drugs according to renal function
- ▶ Diet: sufficient energy and protein while avoiding high protein intake (as renal function is poor, metabolic wastes from protein i.e. urea will accumulate in blood leading to uraemia)
- ▶ **Renal replacement therapy**: if not improving with the above measures

# 4.11 Chronic kidney disease (CKD)<sup>5</sup>

#### Definition

- ► Irreversible loss of renal function developing over years (Davidson)
- ► Abnormalities of kidney structure or function, present for > 3 months, with implications for health (KDIGO)<sup>6</sup>
- ► End-stage renal disease (ESRD): death likely without RRT (CKD stage 5)

# Staging

Stage	Definition	Description	Features
1	$GFR \geq 90  mL/min/1.73m^2$ with evidence of kidney damage	Normal function	Asymptomatic
2	GFR 60-89	Mild CKD	Asymptomatic
3A	GFR 45-59	Mild to moderate CKD	Usually asymptomatic
3B	GFR 30-44	Moderate to severe CKD	Anaemia in some patients
4	GFR 15-29	Severe CKD	First symptoms often at GFR < 20. Electrolyte disorders
5	GFR < 15	Kidney failure	Significant symptoms and complications. Dialysis initiated if < 10.

Table 4.1: Stages of CKD

### **Causes**

- ▶ Diabetes mellitus
- ► Hypertension
- ► Glomerular diseases (IgA nephropathy most common)
- ► Tubulointerstitial diseases
- ► Systemic inflammatory diseases (SLE, vasculitis)
- ► Renal artery stenosis
- ► Congenital / inherited (Alport's, PKD)

#### **Features**

► Typically detected incidentally (by raised urea & creatinine) during routine tests, especially in high-risk patients (e.g. DM, Htn)

<sup>&</sup>lt;sup>5</sup>Here's an awesome video by Ninja Nerd that discusses CKD in detail with its pathophys, features, dx, and

 $<sup>^6</sup> https://kdigo.org/wp-content/uploads/2017/02/KDIGO\_2012\_CKD\_GL.pdf$ 

- ▶ Most cases asymptomatic until GFR  $\geq$  30 mL/min
- ▶ *Nocturia* is an early symptom, due to loss of concentrating ability.
- ▶ When GFR falls to 15-20, symptoms and signs are common.
  - o Tiredness, breathlessness (due to anaemia, fluid overload)
  - Pruritus
  - o Anorexia, nausea, vomiting, hiccups
  - o Wt loss
  - o Kussmaul breathing if profound metabolic acidosis

#### Features according to systems

- ▶ *Bone* (renal osteodystrophy; mechanism Fig. 4.3)
  - o Osteomalacia (renal rickets; due to ↓ vit-D)
  - o Osteoporosis
  - o Osteosclerosis (↑ PTH → rugger jersey spine)
  - o Osteitis fibrosa cystica
- ► Skin
  - o Pruritus
- ► Gastrointestinal
  - o Anorexia
  - o Nausea
  - Vomiting
- ► Metabolic
  - **Hyponatraemia**, **hyperkalaemia** (due to failure of RAAS system)
  - o Hypocalcaemia, hyperphosphataemia (mechanism Fig. 4.3)
  - Metabolic acidosis
  - o Hyperuricaemia, gout
  - o Hypercholesterolaemia
- **►** Endocrine
  - o 2° hyperparathyroidism
  - o Hyperprolactinaemia
- ► Cardiovascular
  - o **Hypertension** ( $\downarrow$ GFR →  $\uparrow$ Renin →  $\uparrow$ Angiotensin)
  - o Heart failure
  - o LVH (2° to anaemia or htn)
  - o Arrhythmias
  - o Pericardial tamponade (uraemic pericarditis)  $\rightarrow \uparrow$  JVP, pulsus paradoxus
- ► Respiratory
  - o Pulmonary oedema

- ► *Nervous*: due to uraemic encephalopathy (azotaemia = elevated urea *without symptoms* i.e. no organ damage yet, uraemia = elevated urea *with symptoms*)
  - o Fatigue, drowsiness
  - Restless leg syndrome
  - o Asterixis (flapping tremor)
  - Seizures
  - o Coma
- ► Muscular

### Investigation

- ▶ Urea and creatinine: to assess stability / progression; compare to previous results
- ► CBC: to see Hb%
- ▶ RBS and HbA1c: to see DM
- ▶ Urinalysis: if haematuria / proteinuria indicate glomerular cause
- ▶ Serum electrolytes: to see Na, K, acidosis
- ightharpoonup Ca, PO $_4^{3-}$ , PTH, vit-D
- ► Lipid profile

### Management

#### Monitoring renal function

- ▶ GFR at least every 6 months for stage  $\geq$  3
- ▶ Plot GFR against time

# Reducing rate of progression

- ► **Antihypertensive** therapy: Slows rate of decline irrespective of the medication
- ► Glycaemic control: if DM present
- **▶** Proteinuria reduction:
  - o Protein
  - $\circ \ \downarrow Proteinuria \rightarrow risk \ of \ progression$ 
    - As protein in the tubules trigger inflammatory damage
  - ACEi / ARBs can both reduce BP and reduce proteinuria
    - $\bullet$  Reduce perfusion pressure by dilating efferent arterioles  $\rightarrow \ \downarrow$  proteinuria
    - Should be prescribed even if no htn
    - Reduce risk of cardiovascular events and all-cause mortality in CKD
    - Angiotensin-II critical for GFR autoregulation in case of low renal perfusion, so may exacerbate, e.g., pre-renal AKI. So should be warned to stop taking the meds if fever, diarrhoea, vomiting (and other potential causes of dehydration / prerenal AKI) arise, and restart after getting better.
    - Should not be commenced if baseline  $K^+ > 5.5$  mmol/L, as they cause hyper-kalaemia

#### **Treatment of complications**

- ► Fluid & electrolyte balance
  - **Limit protein intake**, especially in stages 4 and 5, as urea will accumulate.
  - o Limit potassium intake
  - o Limit salt intake if oedema and htn

#### ► Acid-base balance

- $\circ \downarrow$  excretion of acids  $\rightarrow$  HAGMA (high anion gap metabolic acidosis)
- o NaHCO<sub>3</sub> supplements

#### ► Renal osteodystrophy

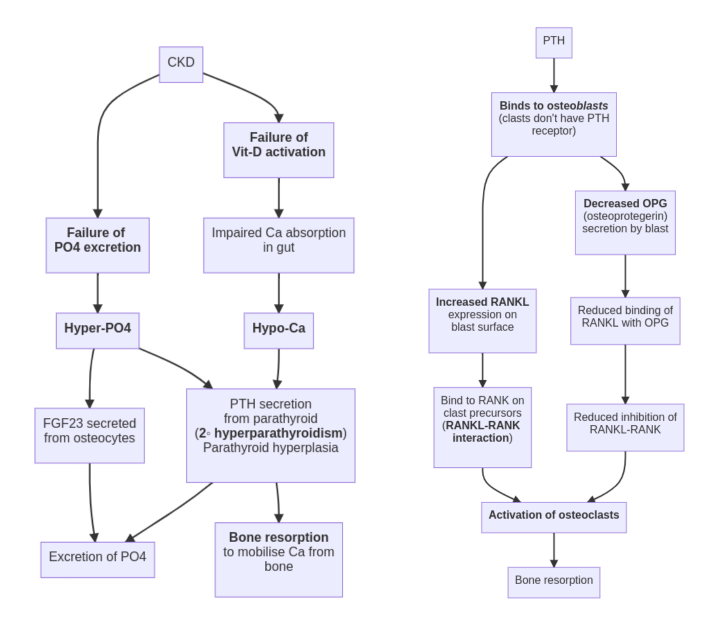
- Primary factors:  $\downarrow PO_4^{3-}$  excretion & vit-D activation (details: Figure 4.3)
- o Phosphate correction:
  - Limit high phosphate food e.g. milk, cheese, eggs, protein-rich food
  - Phosphate-binding drugs e.g. CaCO<sub>3</sub>, Al(OH)<sub>3</sub>
- Vit-D correction:
  - Vit-D supplements

#### ► Anaemia

- o Causes:
  - ↓ erythropoietin
  - Toxic effects of uraemia on RBCs
  - ↓ RBC survival
  - $\uparrow$  capillary fragility  $\rightarrow$  blood loss
  - ullet  $\downarrow$  intake, uptake, and utilisation of dietary Fe
- o Treatment: recombinant human erythropoietin
- o Target Hb value: 10 g/dL

#### ► Cardiovascular risk factors

- Mechanism of ↑ CV disease risk in CKD
  - ↑ BP
  - ullet Vascular tunica media calcification o stiffness
  - Inflammation, oxidative stress
  - Abnormal endothelial function
- o Treatment:
  - Regular exercise
  - Weight loss
  - Smoking cessation
  - Statins, low-dose aspirin



- (a) Mechanism of renal osteodystrophy
- (b) Mechanism of PTH-induced bone resorption

Figure 4.3: PTH and renal osteodystrophy

### 4.12 UTI

#### **Definition**

▶ Presence of >  $10^5$  organisms/mL in a mid-stream sample of urine (if asymptomatic).

#### **Features**

- ► LUTI: cystitis/urethritis
  - Frequency
  - Urgency
  - o Dysuria (burning urethral pain during micturition)
  - o Haematuria
  - Strangury (intense desire to pass more urine after voiding, due to spasm of inflamed bladder wall)
- ▶ UUTI: acute pyelonephritis
  - o Fever with chills and rigor
  - Vomiting
  - o Loin pain
  - o Renal angle tenderness

### Commonly involved pathogens

- ► E. coli: 75%
- ► Proteus
- ► Pseudomonas
- ► Streptococci
- ► Staph. epidermidis

# Investigation

- ▶ Dipstick test for nitrites, leucocyte esterase, and glucose
  - o Most urinary pathogens (e.g. E. coli, Proteus etc) reduce nitrate to nitrite
  - o  $UTI \rightarrow Neutrophils$  in urine  $\rightarrow$  leucocyte esterase
- ▶ Microscopy for WBC and organisms
- ▶ Urine culture

#### **Treatment**

#### **Cystitis**

- ▶ 1st choice
  - o Trimethoprim (200mg bds 3 days)
  - Nitrofurantoin (50mg qds 3 days)
- **▶** Pregnancy

- o Nitrofurantoin (50mg qds 7 days)
- o Cefalexin (250mg qds 7 days)
- Avoid trimethoprim during pregnancy, and nitrofurantoin at term

### **Pyelonephritis**

- ▶ 1st choice
  - o Cefalexin (1g qds 14 days)
  - o Ciprofloxacin (500mg bds 7 days)
- ► Hospitalise if no response within 24h

#### **Epididymo-orchitis**

► 1st choice: Ciprofloxacin

#### Acute prostatitis

▶ 1st choice: Trimethoprim

### Prophylactic measures in women with recurrent UTI

- ► Fluid intake  $\geq 2L/day$
- ► Regular complete bladder evacuation
- ▶ Emptying the bladder before and after intercourse
- ► Good personal hygiene
- ► Continuous prophylactic trimethoprim (100mg) and nitrofurantoin (50 mg) at night

# 4.13 Renal cell cancer

#### Introduction

- ► Peak incidence: 65-75y
- ► Arises from *renal tubular cells*
- ► Histological types:
  - o Clear cell (most common; 85%)
  - o Papillary
  - o Chromophobe
  - Collecting duct

#### **Features**

- ightharpoonup pprox 50% found incidentally (i.e. asymptomatic when diagnosed)
- ightharpoonup pprox 10% present with the **classic triad** (especially if advanced disease)
  - **Haematuria** ( $\approx 60\%$  of symptomatics)
  - $\circ$  Loin pain ( $\approx 40\%$  of symptomatics)

- Loin mass ( $\approx 25\%$  of symptomatics)
- ► Pyrexia of unknown origin (PUO)
- ► Anorexia, malaise, wt loss
- ▶ **Metastasis:** ( $\approx \frac{1}{4}$ th have mets at presentation)
  - Lymphatic: to *para-aortic nodes* Blood-borne: to *lungs, bones, brain*

# Investigation

- ▶ USG: to differentiate solid tumour and cyst
- ► CECT (contrast-enhanced CT) of abdomen and chest: for staging ("cannon-ball" opacities in chest)
- ► CBC: polycythaemia (due to ↑ erythropoietin)
- ► ALP: to check bony mets

### Management

- ► Radical nephrectomy
- ▶ **Partial nephrectomy**: If tumour  $\leq 4$ cm
- ▶ If high operative risk / patient doesn't want surgery:
  - Cryotherapy
  - o Radiofrequency ablation
- ▶ If unresectable / metastatic:
  - o RCC is radio-resistant and also resistant to most chemo agents
  - Current drugs of choice:
    - Tyrosine kinase inhibitors:
      - pazopanib, sunitinib
      - inhibit angiogenesis
    - mTOR inhibitors (mammalian target of rapamycin):
      - temsirolimus, everolimus
  - o High-dose IL-2

# 4.14 Stuff not large enough to devote a section to

# Chapter 5

# Rheumatology

# 5.1 Investigation of musculoskeletal disease

# Joint fluid aspiration

- ► Normal:
  - o Amount small
  - Viscosity high
  - Colourless / pale yellow
- ▶ Inflammation:
  - Amount raised
  - Viscosity lowered (due to enzymatic degradation of hyaluronan & aggrecan)
  - Turbid (due to neutrophils)
- Crystal-induced arthropathies
  - Crystals seen by polarised light microscopy
  - $\circ$  Urate crystals  $\rightarrow$  long, needle shaped, -ve birefringence
  - Ca pyrophosphate crystals → small, rhomboid, +ve birefringence

# Bone scintigraphy

- ▶ Dx of metastatic bone disease and Paget's
- ▶ <sup>99</sup>Tc radiolabelled bisphosphonate used

# **DEXA (Dual Emission X-ray Absorptiometry)**

- ► Measure BMD (bone mineral density)
  - $\circ$  < -2.5  $\rightarrow$  osteoporosis
  - o Between -2.5 and -1  $\rightarrow$  osteopoenia
  - $\circ$  > 2.5  $\rightarrow$  high bone mass (most common cause osteoarthritis)

### **Immunology**

#### ► RF

- o Antibody to Fc fragment of human Ig
- o 70% sensitive for RA (if nodules & extra-articular manifestations then 100% sensitive); specificity poor
- o RF +ve diseases
  - Rheumatoid arthritis
  - Sjogren's syndrome
  - SLE
  - Old age (> 65)

#### ► ACPA

- o Antibody to peptides in which arginine has been converted to citrulline by peptidylarginine deiminase, an enzyme abundant in inflamed synovium.
- o 70% sensitive, >95% specific for RA
- ► ANA (antinuclear antibodies)
  - o 100% sensitive for SLE but poor specificity
  - o ANA +ve diseases
    - SLE
    - Sjogren's
    - Systemic sclerosis
    - Rheumatoid arthritis

### ► Complement C3

• Active SLE  $\rightarrow \downarrow$  C3 (due to consumption of C3 by immune complexes)

# 5.2 Seropositive vs Seronegative arthritis

- ► Seropositive: RF+ inflammatory arthritis
  - o Rheumatoid arthritis
  - o SLE
- ► Seronegative: RF- inflammatory arthritis
  - Ankylosing spondylitis
  - Reactive arthritis
  - Psoriatic arthropathy

# 5.3 Osteoarthritis

- ► Characterised by
  - o degeneration of articular cartilage
  - subchondral osteosclerosis

- o osteophyte formation at joint margin
- o enlargement of affected joint

#### ➤ Sites

- o hips
- o knees
- o PIPs
- o DIPs
- o cervical and lumbar spine

#### ► Investigation:

- X-ray of affected joint: findings described above in characteristics
- o MRI spine if spine OA + suspected root compression / spinal stenosis

#### **▶** Treatment

- o Conservative:
  - Wt loss
  - Exercise
  - NSAIDs
  - Intraarticular glucocorticoids
- o Surgical: if refractory
  - Total joint replacement
  - Osteotomy

# 5.4 Spondyloarthropathies

- ► Asymmetrical oligoarthrites associated with HLA-B27 and typically involving the spine
  - Ankylosing spondylitis
  - Reactive arthritis
  - Psoriatic arthropathy
  - Axial spondyloarthritis
  - Entropathic spondyloarthritis (arthritis associated with IBD)

#### ▶ Common features:

- Asymmetric oligoarthritis
- Sacroilitis
- o Enthesitis (inflammation where tendon attaches to bone)

#### Reactive arthritis

- ▶ "Reactive" to certain infections e.g. Chlamydia, Campylobacter Salmonella, Shigella.
- ► Reiter's syndrome:
  - Triad of can't see, can't pee, can't bend the knee
    - Conjunctivitis

- Urethritis
- Reactive arthritis
- o Due to *Chlamydia*

# Chapter 6

# Neurology

### 6.1 Raised ICP

► Normal ICP = 5-15 mmHg

#### **Causes**

- ► ICSOL
  - Intracranial haemorrhage
  - o Tumours e.g. glioma
  - Brain abscess
- ► **Hydrocephalus**: blockade of CSF circulation
  - Obstructive / non-communicating
  - o Communicating
- ► Cerebral oedema e.g. meningoencephilitis
- ▶ Venous sinus obstruction e.g. cerebral venous thrombosis

#### **Features**

- ▶ Headache
- **▶** Vomiting
- ▶ **Diplopia / blurred vision**: Due to 6th nerve palsy
  - o 6th nerve palsy due to
    - stretching of the long, slender nerve
    - compression against petrous temporal bone
  - o This palsy of the 6th nerve secondary to raised ICP is known as a *false localisation sign*. If the patient presented only with visible features of 6th nerve palsy, (e.g. diplopia, medial squint) we would falsely localise the primary defect to the 6th nerve. So in 6th nerve palsy always exclude RICP by looking for papilloedema.
- **▶** Depressed consciousness
- ► Papilloedema

- **▶** Bradycardia
- **▶** Hypertension

### Management

- ► According to cause:
  - $\circ$  Mass lesion  $\rightarrow$  surgical decompression
  - $\circ$  Hydrocephalus  $\rightarrow$  *ventriculoperitoneal shunt* operation
  - $\circ$  Oedema  $\rightarrow$  glucocorticoids
- ► Supportive:
  - Head elevation
  - o Fluid balance
  - o BP control
  - o Diuretics: mannitol

# 6.2 Neurological emergencies

- **▶** Status epilepticus
- ► Stroke (if thrombo)
- ► Subarachnoid haemorrhage
- **▶** Cord compression
- ► GBS
- ► Myasthenia gravis (if bulbar and/or respiratory)

# 6.3 Status epilepticus

#### **Definition**

- ▶ Continuous or recurrent seizures for  $\geq$  30 mins without gain of consciousness in between.
- ▶ Clinically we assume SE after 5mins of seizure activity.

### Management

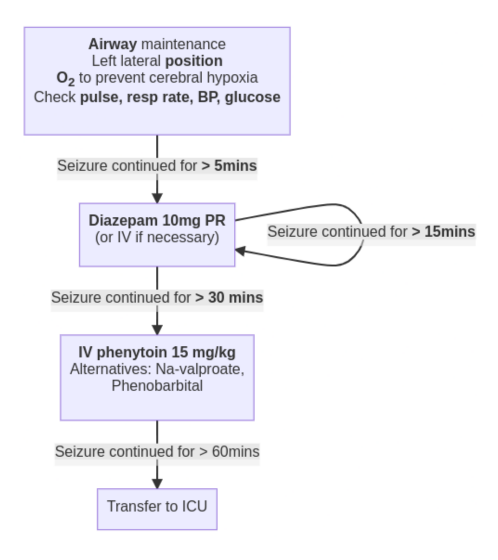


Figure 6.1: Management algorithm for status epilepticus

# 6.4 All jerks root values

Biceps: C5Supinator: C6Triceps: C7

► Finger (aka Hoffmann test): C8

▶ Knee: L3, L4▶ Ankle: S1, S2

▶ Plantar: S1 (technically not a jerk since it's a superficial reflex)

# 6.5 Subarachnoid haemorrhage

#### **Causes**

- ► Ruptured berry aneurysm (85%)
- ► Arterio-venous malformations

#### **Features**

- ► Sudden severe "thunderclap" headache (often occipital)
- ▶ Vomiting
- ► High BP
- ► Neck stiffness
- ► May be loss of consciousness
- ► Photophobia

# Investigation

- ► CT scan: hyperdense material in the subarachnoid space
- ► Lumbar puncture: blood, xanthochromia

# Management

- ▶ Nimodipine 30-60mg IV for 5-14d, followed by 360mg oral for 7d
  - o prevents delayed ischaemia
- ► Insertion of **Pt coils** into aneurysm
- ► **Surgical clipping** of the neck of the aneurysm
- ► Surgical removal if AVM

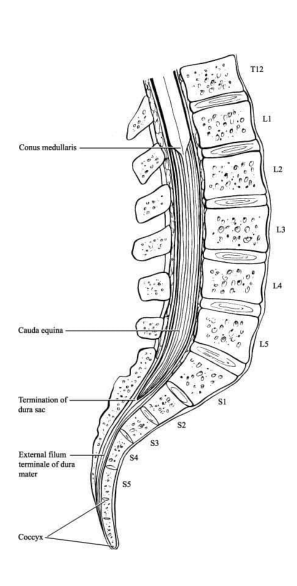
# 6.6 Subacute combined degeneration

#### **Features**

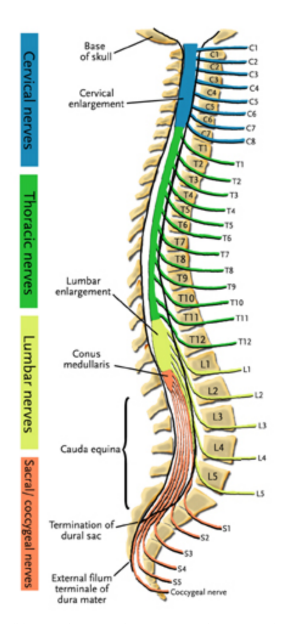
- ▶ Peripheral neuropathy: due to demyelination
- ► Signs of dorsal column lesion: position and vibration sense lost
- ► Signs of **pyramidal lesion**:
  - o plantar extensor
  - o knee jerk brisk
  - ankle jerk absent: as peripheral neuropathy affects longer nerves first, the afferent pathway for ankle jerk is damaged by the PN while the same for knee jerk is not. Combined with the corticospinal tract lesion, this makes knee jerks brisk and ankle jerks absent.
- ▶ Optic atrophy: death of retinal ganglion cell axons

# 6.7 Cauda equina and Conus medullaris lesions

# **Anatomy**



(a) The Conus and the Cauda



(b) Vertebrae with corresponding spinal cord segments

- ▶ During development, the spine grows faster than the spinal cord, which is why the spinal nerves exit the spinal column at increasingly oblique angles.
- ► The spinal cord ends as the tapered conus medullaris at around L1.

# Relevant physiology

- ► S2-4:
  - o Parasympathetic fibres for bladder sphincter (activation causes emptying)
  - $\circ$  Somatic fibres for pudendal nerves (activation causes relaxation of urethral and anal sphincters  $\rightarrow$  emptying).
- ► Conus is surrounded by spinal nerve roots bundled up together around it, so injury / compression in this region leads to a combination of UMN and LMN lesion features.
- ▶ Lesion in the cauda region leads only to LMN lesion features as there's no cord there.

# Features of Cauda Equina syndrome vs Conus Medullaris syndrome

Features	Cauda Equina Syndrome	Conus Medullaris
Vertebral level	L2-sacrum	L1-L2
Spinal level	Injury to the lumbosacral nerve roots	Injury of the sacral cord segment (conus and epiconus) and roots
Severity of symptoms and signs	Usually severe	Usually not severe
Symmetry of symptoms and signs	Usually asymmetric	Usually symmetric
Pain	Prominent, asymmetric, and radicular	Usually bilateral and in the perineal area
Motor	Weakness to flaccid paralysis	Normal motor function to mild or moderate weakness
Sensory	Saddle anesthesia, may be asymmetric	Symmetric saddle distribution, sensory loss of pin prick, and temperature sensations (Tactile sensation is spared.)
Reflexes	Areflexic lower extremities; bulbocavernosus reflex is absent in low CE (sacral) lesions	Areflexic lower extremities  (If the epiconus is involved, patellar reflex may be absent, whereas bulbocavernosus reflex may be spared.)

Reflexes	Areflexic lower extremities; bulbocavernosus reflex is absent in low CE (sacral) lesions	Areflexic lower extremities  (If the epiconus is involved, patellar reflex may be absent, whereas bulbocavernosus reflex may be spared.)
Sphincter and sexual function	Usually late and of lesser magnitude; lower sacral roots involvement can cause bladder, bowel, and sexual dysfunction	Early and severe bowel, bladder, and sexual dysfunction that results in a reflexic bowel and bladder with impaired erection in males
EMG	Multiple root level involvement; sphincters may also be involved	Mostly normal lower extremity with external anal sphincter involvement
Outcome	May be favorable compared with conus medullaris syndrome	The outcome may be less favorable than in patients with CES

# 6.8 Neurogenic Bladder

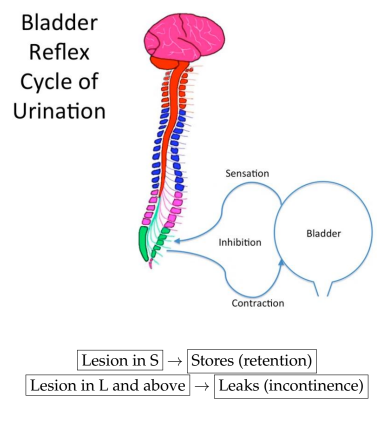
# Physiological control of micturition: the Micturition Reflex

"Components"

- **▶** Detrusor muscle
  - Muscles of the bladder wall
  - o Innervated by
    - Parasympathetic:
      - cholinergic M<sub>3</sub> receptors
      - pelvic nerve from **S2-4**
      - $\blacksquare$  causes contraction  $\rightarrow$  urination
    - Sympathetic:
      - $\blacksquare$   $\beta_3$  receptors
      - hypogastric nerve from **T10-L2**
      - $\blacksquare$  causes **relaxation**  $\rightarrow$  **storage** / retention
- ► Internal urethral sphincter
  - o Innervated by only sympathetic
    - hypogastric nerve from T10-L2

- causes storage / retention
- **►** External urethral sphincter
  - Voluntary control
    - UMN from pontine micturition centre
    - LMN from S2-4 (pudendal nerve)
    - causes retention

#### The big picture (highest yield for clinical interpretation)



- ▶ The reflex spontaneously tries to void the bladder upon being filled even by small amounts. The only thing holding it back is the pontine micturition centre which always inhibits the circuit, unless we voluntarily signal it not to. We can't signal the bladder to empty itself. We can only signal the micturition reflex to not hold back.
- ➤ The reflex circuit is composed of afferents and efferents from S2-4. So any **lesion above S2-4** will leave the circuit intact, while damaging the telephone lines from the pontine centre. The end result is that we lose the inhibition of the reflex (which fires every few mins after storing some tens of millilitres of urine), and without anything holding it back, there's urinary **incontinence**.
- ▶ On the other hand, any **lesion at S2-4** will damage the circuit elements themselves. That means that the reflex will be lost. This means we no longer have access to the bladder emptying circuit. So there will be urinary **retention**.

#### The details

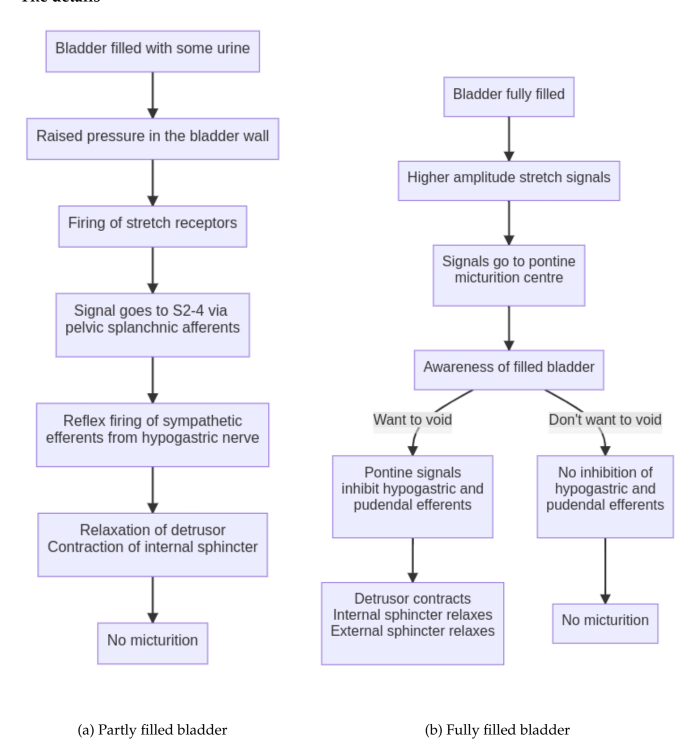


Figure 6.3: Neural control of micturition

## 6.9 Parkinson's disease

### **Parkinsonism**

- ► Triad of TRH: tremor, rigidity, hypokinesia (bradykinesia)
- **▶** Causes
  - Idiopathic → Parkinson's disease
  - o Cerebrovascular disease
  - o Drugs:
    - Antipsychotics (older)
    - Metoclopramide ( $D_2$  blocker  $\rightarrow$  undo  $D_2$  mediated inhibition of cholinergic stimulation in GIT  $\rightarrow \uparrow$ GI motility  $\rightarrow$  antiemesis)
      - Domperidone, which is also a  $D_2$  blocker, selectively acts on  $D_2$ s in the GIT, so no parkinsonism
      - Metoclopramide is nonselective
    - Na-valproate, Lithium
  - o Other neurodegenerative diseases
    - Lewy body dementia
    - Multiple system atrophy
    - Alzheimer's
  - o Genetic
    - Huntington's
    - Wilson's

## Pathophysiology

- ► Loss of pigmented dopaminergic neurons in substantia nigra
- ► Lewy bodies in nigral cells: pathological hallmark
  - o Eosinophilic cytoplasmic inclusions
  - o Lewy body = aggregation of  $\alpha$ -synuclein (hence the alternative name  $\alpha$ -synucleinopathy)
  - o Other  $\alpha$ -synucleinopathies (diseases associated with Lewy body deposition (differ in the initial site of deposition and hence initial features)):
    - Lewy body dementia
    - Multiple system atrophy

### **Features**

- ► Average age of onset: 60y
- ► First degreee relative with PD  $\rightarrow$  2-3x risk
- ► Progressive, incurable
- ▶ Initially motor symptoms dominate, but eventually nonmotor symptoms (e.g. depression, anxiety, cognitive impairment) become increasingly prominent
- ► Motor symptoms initially asymmetrical
- ► The hallmark is **bradykinesia**, which leads to

- o Micrographia (small handwriting)
- Difficulty tying shoelaces / buttoning clothes
- Difficulty rolling over in bed
- ▶ **Resting tremor** affecting limbs, chin and jaw but not the head
- ► Rigidity
- ► Soft, indistinct speech

## Signs

- ► General:
  - o Hypomimia (expressionless / mask-like face)
  - o Dysphonia (soft, indistinct speech)
  - o Flexed (stooped) posture with impaired postural reflexes
  - o Bradykinesia
  - Glabellar tap:
    - tapping on glabella (above bridge of nose) → blink
    - normal → blinking stops after 3-5 times
    - Parkinson's → sustained blinking
- ► Gait: Festinating gait
  - Slow to start
  - o Short, shuffling steps (festination)
  - o Reduced arm swing
  - o Impaired balance on turning (fractionated turn)
- ► Tremor:
  - First in arm/hand (pill-rolling tremor)
- ► Rigidity:
  - Leadpipe rigidity:
    - better seen in elbow / knee
    - uniform throughout movement
  - o Cogwheel rigidity (= tremor superimposed on rigidity) better seen in wrist
  - o Rigidity vs Spasticity:
    - Rigidity:
      - uniform resistance throughout range of motion
      - due to *extrapyramidal* lesion
    - Spasticity:
      - initially increased resistance, followed by lessening
      - due to *pyramidal* lesion
- ▶ The following will be normal (if abnormal, consider other causes)
  - o Power, jerks, plantar
  - Eye movements
  - Sensory exam
  - Cerebellar exam

## Investigation

- ▶ Dx is clinical
- ► CT / MRI normal
- ▶ Functional dopaminergic imaging (SPECT / PET) abnormal even early
- ▶ In younger, exclude Huntington's and Wilson's

### **Treatment**

- ► Only symptomatic, no cure
- ▶ **Physiotherapy**  $\pm$  **drugs** (drugs not given if mild)
- ► Surgical: on failure of medical therapy

### **Drugs**

- ► Levodopa + carbidopa
  - Most effective in reducing rigidity and bradykinesia
  - **Role of dietary protein**: amino acids in dietary protein compete with levodopa for intestinal absorption and transport across BBB → ↓ efficacy of levodopa
  - Side effects:
    - Postural hypotension
    - Nausea, vomiting
    - Hallucinations
  - o Fluctuating response after 3-5y of use
    - *End of dose dyskinesia*: due to progressive loss of dopamine, duration of action of levodopa becomes shorter. Freezing and rigidity before next dose of levodopa.
      - Management: smaller, frequent dosage
    - *On-off phenomenon*: periods of severe parkinsonism (freezing and immobility  *off period*) alternating with periods of dopamine-induced dyskinesia / chorea (*on period*).
      - Management: lower levodopa dose, add selegiline with levodopa
  - Contraindications:
    - Psychosis
    - Narrow angle glaucoma
    - Malignant melanoma
    - PUD

## Anticholinergics

- o Benzhexol, benztropine, trihexyphenidyl
- o Reduce tremor and rigidity, not bradykinesia
- o Side effects:
  - Urinary retention, constipation, dry mouth
  - Worsening of glaucoma
- o Contraindications:
  - BEP

• Narrow angle glaucoma

## **▶** Dopamine receptor agonists

- Ergot-derived (e.g. bromocriptine) or non-ergot-derived (e.g. ropinirole)
- o Compared to levodopa, less therapeutic effect, more side effects
- o Side effects:
  - Postural hypotension
  - Nausea, vomiting
  - Hallucination
  - Confusion

#### **►** MAO-B inhibitors

- o Selegiline, rasagiline
- o MAO-B facilitates dopamine breakdown → MAO-Bi potentiates the action of levodopa by inhibiting breakdown

#### **►** COMT inhibitors

- o Tolcapone, entacapone
- $\circ$  COMT  $\rightarrow$  peripheral breakdown of levodopa

#### ► Amantadine:

- o Rarely used unless patient unable to tolerate other drugs.
- o Mild, short-lived effect.

## Surgery

### ► Deep brain stimulation

- Replaced destructive surgery
- o Targets: thalamus, globus pallidus, subthalamic nuclei
- $\circ$  Thalamic stimulation  $\rightarrow$  reduce tremor

# **Diabetes Mellitus**

## 7.1 Mechanism of insulin secretion



Figure 7.1: Mechanism of insulin secretion

## 7.2 Incretin effect

For the same glucose load applied orally and IV, the **oral load stimulates more insulin secretion** (because oral load  $\rightarrow$  release of gut peptides GLP-1 and GIP  $\rightarrow \uparrow$  insulin secretion).

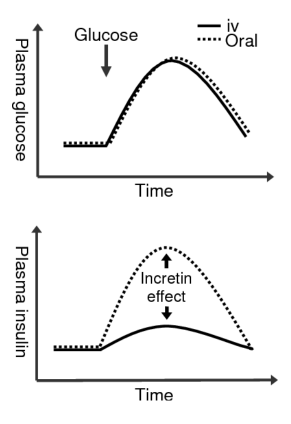


Figure 7.2: The incretin effect

# 7.3 Diagnostic criteria

- ▶ If *symptomatic*, any one of the following
  - $\circ \ FBS \geq 7.0 \ mmol/L$
  - $\circ$  RBS / 2h after OGTT  $\geq$  11.1 mmol/L
  - $\circ$  HbA1c > 6.5%
- ▶ If *asymptomatic*, two positives needed for dx.
- ► IFG:
  - o FBS between 6.1-6.9 mmol/L, 2h after OGTT < 7.8 mmol/L (WHO)
  - o FBS between 5.6-6.9 mmol/L (American diabetes association (ADA))
- ► IGT:
  - o FBS < 7.0 mmol/L, 2h after 7.8-11 mmol/L

## 7.4 Diabetic ketoacidosis (DKA)

- ► Medical emergency
- ► Cause of death
  - o Children: cerebral oedema
  - o Adults:
    - Hypokalaemia
    - ARDS
    - Comorbidities: acute MI, sepsis, pneumonia

#### ► Cardinal biochemical features

- $\hspace{1cm} \circ \hspace{1cm} Hyperglycaemia \rightarrow osmotic \hspace{1cm} diures is \rightarrow dehydration, \hspace{1cm} dyselectrolytaemia \\$
- o Hyperketonaemia:
  - ullet Insulin deficiency + elevated catecholamines o unrestrained lipolysis to make FFA o hepatic ketogenesis
- Metabolic acidosis

### Clinical features

- **▶** Symptoms
  - o Polyuria, thirst
  - o Weakness
  - o Nausea, vomiting
  - o Abdominal pain
  - o Blurred vision

### ▶ Signs

- Dehydration
- Hypotension
- o Tachycardia
- o Air hunger / Kussmaul breathing (deep and sighing breathing)
- o Acetone breath
- o Delirium, drowsiness, coma

## Management

- **▶** Establish IV access
- ▶ Volume replacement: 0.9% NaCl
  - o If systolic BP  $\geq$  90mmHg: 1L over 1h
  - ∘ Else:  $\frac{1}{2}$ L over 15mins → reassess. If BP still < 90mmHg, repeat.
- ► Insulin therapy: IV 0.1 U/kg/h
  - o Corrects hyperglycaemia & acidosis
- **▶** Monitor

- o Every 1h:
  - capillary blood glucose and ketone
- Every 2h: Venous HCO<sub>3</sub> and K<sup>+</sup>
- o Every 4h: Serum electrolytes
- ► If K<sup>+</sup> is low, 40mmol/L KCl with normal saline

# 7.5 Hypoglycaemia

### **Features**

- ► Autonomic
  - Sweating
  - o Trembling
  - o Palpitations

## ► Neuroglycopoenic

- o Delirium
- o Drowsiness
- Speech difficulty
- Incoordination

## Management

- ▶ Oral fast-acting carbohydrate (10-15g) e.g. glucose drink / confectionery / honey to buccal mucosa
- ▶ Repeat capillary glucose measurement 10-15mins later
  - o If still hypo, repeat upto 3 cycles
  - o Still hypo after 3 cycles → glucagon 1mg IM

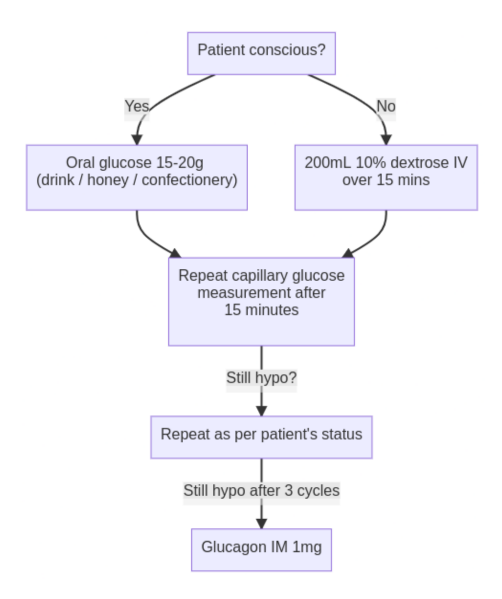


Figure 7.3: Managing hypoglycaemia

# 7.6 Insulin therapy

## **Indications**

- ► Type I DM
- ► Type II DM not controlled by OHA
- ▶ DIP / GDM
- ► DKA
- ► Hyperkalaemia

# **Preparations**

- ► **Rapid-acting** (rapid=LAG-less)
  - o Lispro

- o Aspart
- Glulisine
- ► **Short-acting:** soluble/regular insulin
- ► Intermediate-acting: Isophane (I for I)
- ► Long-acting
  - o Glargine (gLARGE-in)
  - o Detemir Route of administration: subcutaneous

## 7.7 Oral Hypoglycaemic Agents

- **▶ Biguanides**: Metformin
  - o Insulin sensitiser
  - o Mechanism of action
    - \( \text{hepatic glucose production (gluconeogenesis and glycogenolysis)} \)
    - ↑ gut glucose uptake & utilisation
    - weak inhibitor of mitochondrial respiration  $\rightarrow \uparrow$  AMP,  $\downarrow$  ATP  $\rightarrow \uparrow$  glucose uptake utilisation etc.
  - Side effects profile
    - Weight neutral
    - Non-hypoglycaemic
    - Lactic acidosis
- ▶ Sulphonylureas: Glibenclamide, Gliclazide, Glimepiride
  - o Insulin *secretagogue*
  - **Mechanism of action:** Block K<sup>+</sup> channel in  $\beta$ -cells  $\rightarrow \uparrow$  insulin secretion
  - Side effects profile
    - Wt gain
    - Hypoglycaemia
- $ightharpoonup \alpha$ -glucosidase inhibitors: Acarbose
  - Mechanism of action: delay absorption of carbs
  - Side effects profile
    - Non-hypoglycaemic
    - Flatulence
    - Bloating
    - Diarrhoea
- ► Incretin-based therapies:
  - o **DPP-4 inhibitors:** Gliptins
    - MoA
      - DPP-4: breaks down GLP-1 & GIP → inhibit incretin effect
  - o GLP-1 receptor agonists: Exenatide, liraglutide
- ► Thiazolidinediones: Pioglitazone

- Mechanism of action
  - PPAR- $\gamma$  agonist  $\rightarrow$  enhance action of insulin
- Side effects profile
  - Non-hypoglycaemic
  - Wt gain (increase fat cells)
- ▶ SGLT-2 inhibitors: empagliflozin, dapagliflozin
  - o **MoA:** inhibit reabsorption of glucose in renal tubules  $\rightarrow$  25% of filtered glucose excreted
  - o Resulting glycosuria can lead to genital fungal infections
  - o Empagliflozin  $\rightarrow$  35% reduced mortality in heart failure

## 7.8 Complications of DM

## **Acute complications**

- ► Hypoglycaemia
- ► Diabetic ketoacidosis
- ► Hyperglycaemic hyperosmolar state (HHS)
- ► Lactic acidosis

## Long-term complications

### Microvascular

- ▶ Diabetic **neuropathy**: peripheral neuropathy (sensory, motor or mixed), mononeuritis multiplex, autonomic neuropathy
- ► Diabetic **nephropathy**
- ► Ocular complications:
  - Diabetic retinopathy
  - o Cataract
- ► Foot complications: ulcer, gangrene
  - o Causes of diabetic ulcer:
    - ischaemia
    - neuropathy
    - combined ischaemia and neuropathy
    - trauma
    - infection

#### Macrovascular

- ► Coronary: myocardial ischaemia, infarction
- ► Cerebral: Cerebrovascular disease
- ▶ Peripheral: ischaemia, claudication

## 7.9 Pathogenesis of chronic complications

► Thickened basement membrane and ↑ vascular permeability are the pathophysiological hallmarks

### **Mechanisms**

- ► Formation of advanced glycation end-products (AGEs)
  - Nonenzymatic addition of glucose to proteins
  - o Exert effects by both binding to receptors (RAGE) on endothelial, inflammatory and smooth muscle cells, and by direct (non-receptor) cross-linking of collagen in the basement membrane, mediated chiefly by **VEGF** and **TGF-**β (details: fig. 7.4)
- ► Excess activation of **protein kinase C (PKC)** 
  - o Mechanism:  $\uparrow$ intracellular glucose  $\rightarrow \uparrow$ DAG synthesis (diacyl glycerol)  $\rightarrow \uparrow \oplus$ PKC  $\rightarrow \uparrow$ VEGF, TGF- $\beta$ , PAI-1 (plasminogen activator inhibitor-1) production by the vascular endothelium
- ▶ Oxidative stress and sorbitol accumulation
  - Mechanism: ↑intracellular glucose → ↑metabolism to sorbitol (by aldose reductase)
     → fructose using NADPH → NADPH used up → ↓availability for use in antioxidant pathway → ↑ROS-mediated damage
  - $\circ$  Accumulation of sorbitol in the lens  $\rightarrow$  cataract

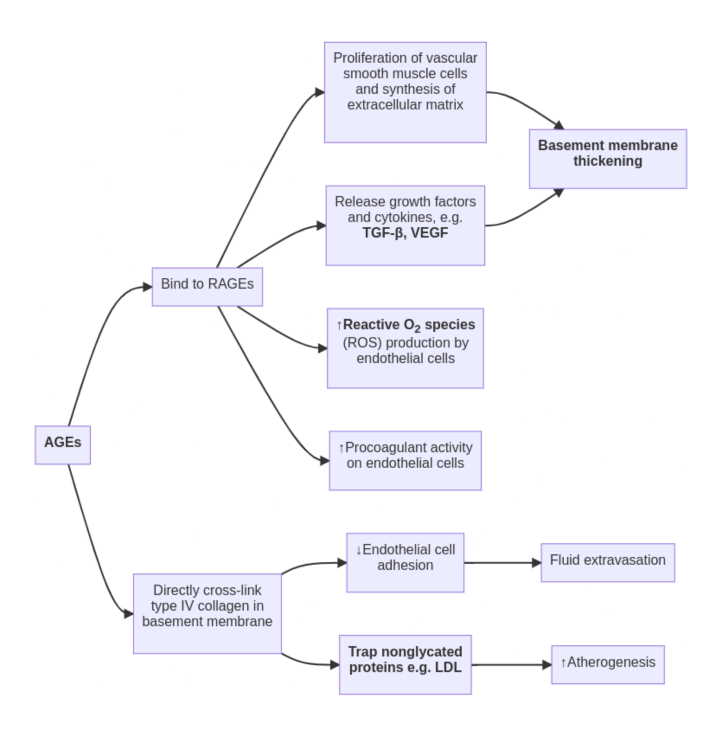


Figure 7.4: Mechanisms of AGE-induced microvascular pathology

# 7.10 Stuff not large enough to devote a section to

## Metabolic syndrome

- ► AKA insulin resistance syndrome or syndrome X
- ► Type 2 DM associated with central obesity + htn + dyslipidaemia (high TGs/LDLs and/or low HDL)
- ▶ Often associated with
  - o Nonalcoholic fatty liver disease (NAFLD)
  - o PCOS
- ▶ ↑risk of stroke, CAD

# Gastrointestinal diseases

# 8.1 Weight loss

### Causes

- **▶** Endocrine
  - o DM (more in type I)
  - Thyrotoxicosis
  - o Addison's
- ▶ GI
  - o Any cause of dysphagia e.g.
    - Stroke
    - MS
    - Ca oesophagus
    - Achalasia cardia
    - Plummer-Vinson syndrome (oesophageal webs+IDA)
  - o Malabsorption syndrome
    - IBD
    - Chronic pancreatitis (due to enzyme insufficiency)
    - Coeliac disease
- **▶** Malignancies
- **▶** Chronic infection
  - o TB
  - o AIDS
- **▶** Psychological
  - Depression
  - o Anorexia nervosa
  - o Bulimia nervosa
  - o Alcoholism

# Haematology

# 9.1 Chronic myeloid leukaemia (CML)

Defining characteristic: Philadelphia chromosome

- ▶ Shortened chr22 by reciprocal translocation with chr9
- ► Results in BCR-ABL fusion gene
- ▶ BCR-ABL codes for a **tyrosine kinase** which influences cell proliferation and survival

#### **Features**

- ► Wt loss
- ► Lethargy
- ► Abdominal discomfort
- ► Splenomegaly
- ▶ Hepatomegaly

#### **Phases**

- ► Chronic
- ► Accelerated
- ▶ Blastic crisis

## Investigation

- ► CBC: anaemia, leucocytosis
- **▶ PBF**:
  - Full range of granulocytic precursors, from *myeloblasts* to *mature neutrophils*.
  - Predominant: neutrophils and myelocytes.
  - Myeloblasts < 10%.

#### **▶** Bone marrow examination:

- Hypercellular marrow
- $\circ \uparrow M/E$  ratio

- ↓ erythrpoiesis
- o ↑ leucopoiesis
- ► Chromosome analysis to detect Ph chromosome

## Management

## Chronic phase

- ▶ 1st line: Tyrosine kinase inhibitors (TKIs):
  - o Imatinib
  - o Dasatinib
  - o Nilotinib
  - o normalise blood count within a month, complete cytogenetic response (disappearance of Ph chr) within 6 months in 90% patients. Resample bone marrow at 6mo to confirm. Thereafter monitor 3-monthly by RT-PCR for BCR-ABL mRNA transcripts.
- ► Allogeneic HSC transplant: if TKI fails
- **▶** Hydroxycarbamide
- ► **Interferon:** in pregnancy

# **Nutritional diseases**

## 10.1 Vitamins

## **B**<sub>1</sub> (thiamin) deficiency

#### **Functions of thiamin**

- ► Cofactor in different pathways of aerobic metabolism of glucose
  - o decarboxylation of pyruvate to acetyl-coA (so bridge between glycolysis and Krebs)
  - $\circ$  decarboxylation of  $\alpha$ -ketoglutarate to succinate in Krebs

#### **Features**

- ► For its pivotal role in aerobic glucose meta, the largest blow is dealt to the tissues most dependent on glucose the **brain** and the **heart**.
- ▶ Beri-beri
  - o Dry: Neurological beri-beri
    - Peripheral neuropathy
    - Wrist/foot drop
    - Korsakoff's psychosis
    - Wernicke's encephalopathy
  - o Wet: Cardiac beri-beri (wet as it causes generalised oedema)
    - Biventricular failure
    - Generalised oedema
    - Pulmonary oedema

#### **Treatment**

- ▶ Wet and Wernicke's should be treated by IV vit-B and C mixture
- ► Korsakoff's: irreversible, nonresponsive to thiamin

## **B**<sub>12</sub> deficiency

## Functions of $B_{12}$

- ► Recycles folate → essential for **cell division** (especially in RBC)
- **▶** Myelination

#### **Features**

- ► Megaloblastic anaemia
- **▶** Glossitis
- ► Neurologic features
  - o Peripheral neuropathy
  - o Autonomic neuropathy
  - o Optic atropy
  - o Subacute combined degeneration of spinal cord

### **Treatment**

- ightharpoonup Vit-B<sub>12</sub> IM
- ▶ If combined folate and  $B_{12}$  deficiency, only folate should not be given without  $B_{12}$ , as  $B_{12}$  gets used up in folate recycling, deteriorating the  $B_{12}$  deficiency symptoms. Always give both together.