Medicine

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Chapter 1

Cardiology

1.1 Presenting problems in CVS disease

Features of benign murmur

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

1.2 ECG

Anatomy of an ECG



1.2. ECG 3

Abnormalities of components

Pathological Q

- Depth > 2mm
- Height > 1mm
- Present in > 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
 - 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: V₁, V₂
 Anterior: V₃, V₄

Lateral: I, aVL, V₅, V₆
Extensive anterior: V₁-V₆
Anterolateral: I, aVL, V₁-V₆

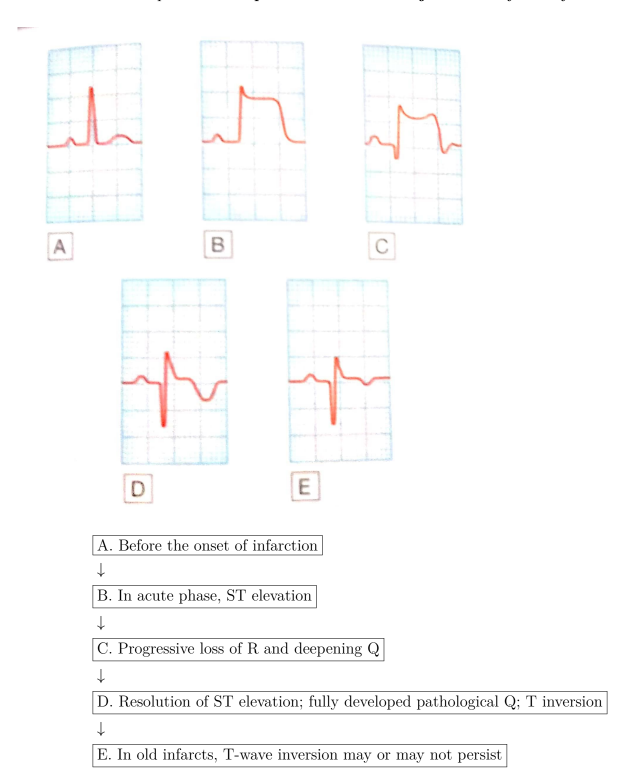
Reciprocal changes

• Acute STEMI in some surface of the heart \to ST elevation in corresponding leads, and ST depression in reciprocal leads

Site	Facing	Reciprocal
Septal	V1, V2	V7, V8, V9
Anterior	V3, V4	None
Lateral	I, aVL, V5, V6	II, III, aVF
Inferior	II, III, aVF	I, aVL
Posterior	V7, V8, V9	V1, V2

Basic pathophys of STEMI

• Occurs due to proximal complete occlusion of major coronary artery



1.2. ECG 5

NSTEMI

• Partial occlusion of major or complete occlusion of minor coronary artery

- Subendocardial/partial-thickness $MI \to \mathbf{no}$ pathological \mathbf{Q}
- ullet ST depression + T inversion in chest leads

Chapter 2

Dermatology

2.1 Anatomy and physiology

- Layers of skin:
 - Epidermis: further layered into (from out→in)
 - * corneum
 - * lucidum
 - * granulosum
 - * spinosum
 - * basale
 - Dermis: contains
 - * blood vessels
 - * nerves
 - * pilosebaceous units (hair follicle + sebaceous gland)
 - Subcutis: adipose

Epidermal appendages

- Hair follicles:
 - 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
 - usually associated with a hair follicle

$$\begin{array}{c} - \ \, \boxed{androgens} \to \uparrow \ \, \boxed{sebum} \\ - \ \, \boxed{oestrogen} \to \downarrow \ \, \boxed{sebum} \end{array}$$

- Sweat glands
 - innerved by sympathetic cholinergic fibres

2.2 Principles of management of skin disease

Topical treatments

- Ointments vs Creams
 - 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
 - Moisturise, lubricate, protect skin
 - Vehicles without active drug
- Gluocorticoids

Phototherapy

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

Systemics

- Antihistamines
- Retinoids
 - Anti-inflammatory
 - Promote differentiation of skin cells

- Teratogenic

- * must be prescribed with robust contraception
- * females must have negative pregnancy test before, during, and after therapy

• Immunosuppressants

- Glucocorticoids e.g. prednisolone
- Methotrexate
- Azathioprine

Biologics

- Biological inhibitors of proinflammatory cytokines
- TNF- α inhibitors
 - Infliximab
 - Etanercept
- Interleukin inhibitors
 - Ustekinumab: IL-12, 23
 - Guselkumab: IL-23Secukinumab: IL-17
- Rituximab:
 - Binds to CD20 \rightarrow cause ADCC of B cells
 - As terminally differentiated plasma cells don't have CD20 they're safe
 - Use: pemphigus vulgaris

Non-surgical therapy

- Cryo
 - Liquid N₂
 - Causes cell membrane destruction \rightarrow death
- Laser
- PDT / photodynamic therapy

2.3 Skin cancers

Classification

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

2.4 Fungal infections

Types

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

2.5 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
 - 2 applications
 - 1 wk apart
 - Whole body, except head
- Oral Ivermectin:
 - Single dose
 - For poor adherence, immunosuppresion or heavy infestation

2.6 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
 - Topical Benzoyl peroxide
 - Topical Retinoids
 - Topical antibiotics
 - * Erythromycin
 - * Clindamycin
- Moderate disease: topical plus
 - Systemic tetracycline
 - Oestrogen containing OCP
 - Isotretinoin: if inadequate response to topical+systemic therapy for 6 months
- Severe disease
 - 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

2.7. ECZEMAS 11

2.7 Eczemas

• Seborrhoeic dermatitis is associated with Malassezia yeasts

Features

Most types have the following clinical features:

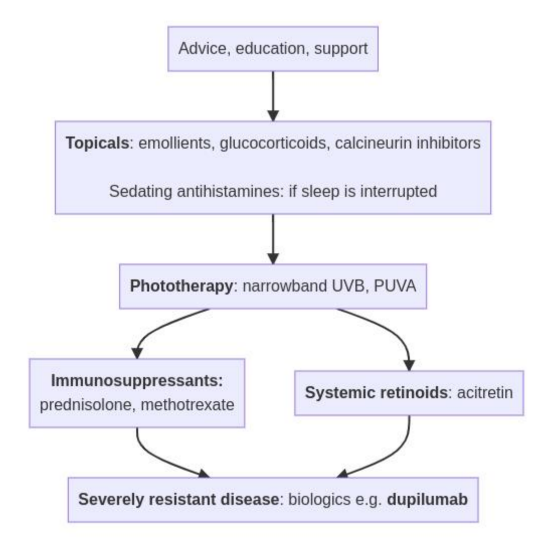
Acute

- Ill-defined erythema, oedema
- Papules, vesicles, bullae
- Exudation
- Scaling

Chronic

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

Management of eczema



2.8. PSORIASIS

2.8 Psoriasis

- Chronic inflammatory hyperproliferative skin disease
- Characteristics
 - Well-defined erythematous scaly plaques
 - 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
- Trauma
- Infection
 - $-\beta$ -haemolytic strep \uparrow guttate psoriasis
 - HIV may initally present with severe psoriasis
- Drugs
 - Antimalarials
 - $-\beta$ -blockers
 - Lithium
 - NSAIDs
- Stress and anxiety

Clinical types

- Plaque psoriasis:
 - most common
 - well-demarcated erythematous plaques
 - silver-white scales in untreated
 - * bleed on scraping (due to dilated vessels underneath) \rightarrow Auspitz sign
 - Sites
 - * extensor surfaces
 - · elbows
 - · knees
 - · lower back
 - * scalp
 - * nails

- Guttate psoriasis:
 - follows Strep throat
 - common in children/adolescent
 - UVB highly effective
 - $-\,$ may he rald the onset of plaque psoriasis in a dulthood
- Erythrodermic sporiasis: generalised \rightarrow medical emergency
- Pustular psoriasis

2.8. PSORIASIS

Management of psoriasis



2.9 Hypopigmentation

Causes

- Vitiligo
- Albinism
- Pityriasis alba
- Pityriasis versicolor

Vitiligo

- Acquired
- Cell-mediated autoimmune destruction of melanocytes
- Loss of melanocytes \rightarrow hypopigmented patches

Albinism

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

2.10 Hyperpigmentation

Causes

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

• Drugs

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

2.11 Pseudorandom factoids

SPF (sun protection factor)

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

Mechanism of venous ulceration



Chapter 3

Neurology

3.1 Raised ICP

• Normal ICP = 5-15 mmHg

Causes

- ICSOL
 - Intracranial haemorrhage
 - Tumours e.g. glioma
 - Brain abscess
- Hydrocephalus: blockade of CSF circulation
 - Obstructive / non-communicating
 - 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
 - 6th nerve palsy due to
 - * stretching of the long, slender nerve
 - * compression against petrous temporal bone
- Papilloedema
- Bradycardia
- Hypertension
- Depressed consciousness

Management

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

3.2 Neurological emergencies

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