Dermatology

A handbook for medical students & junior doctors



Derr	natology: Handbook for medical students & junior doctors
This publication is supported by the	e British Association of Dermatologists.

First edition 2009

Revised first edition 2009
Second edition 2014

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Preface

This Handbook of Dermatology is intended for senior medical students and newly qualified

doctors.

For many reasons, including modern medical curriculum structure and a lack of suitable

patients to provide adequate clinical material, most UK medical schools provide inadequate

exposure to the specialty for the undergraduate. A basic readable and understandable text

with illustrations has become a necessity.

This text is available online and in print and should become essential reading. Dr Chiang is to

be congratulated for her exceptional industry and enthusiasm in converting an idea into a

reality.

Julian Verboy

Professor of Dermatology

Liverpool 2009

Preface to the 2nd edition

Nicole and I are gratifed by the response to this Handbook which clearly fulfils its purpose.

The positive feedback we have received has encouraged us to slightly expand the text and

allowed us to update where necessary. I should like to thank the BAD for its continued

support.

Julian Verbov

Professor of Dermatology

Liverpool 2014

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Foreword to First edition

There is a real need for appropriate information to meet the educational needs of doctors at all levels. The hard work of those who produce the curricula on which teaching is based can be undermined if the available teaching and learning materials are not of a standard that matches the developed content. I am delighted to associate the BAD with this excellent handbook, designed and developed by the very people at whom it is aimed, and matching the medical student and junior doctor curriculum directly. Any handbook must meet the challenges of being comprehensive, but brief, well illustrated, and focused to clinical presentations as well as disease groups. This book does just that, and is accessible and easily used. It may be read straight through, or dipped into for specific clinical problems. It has valuable sections on clinical method, and useful tips on practical procedures. It should find a home in the pocket of students and doctors in training, and will be rapidly worn out. I wish it had been available when I was in need, I am sure that you will all use it well in the pursuit of excellent clinical dermatology!

Dr Mark Goodfield

President of the British Association of Dermatologists

What is dermatology?

 Dermatology is the study of both normal and abnormal skin and associated structures such as hair, nails, and oral and genital mucous membranes.

Why is dermatology important?

- Skin diseases are very common, affecting up to a third of the population at any one time.
- Skin diseases have serious impacts on life. They can cause physical damage, embarrassment, and social and occupational restrictions. Chronic skin diseases may cause financial constraints with repeated sick leave. Some skin conditions can be life-threatening.
- In 2006-07, the total NHS health expenditure for skin diseases was estimated to be around £97 million (approximately 2% of the total NHS health expenditure).

What is this handbook about?

- The British Association of Dermatologists outlined the essential and important learning outcomes that should be achieved by all medical undergraduates for the competent assessment of patients presenting with skin disorders (available on: http://www.bad.org.uk/Portals/_Bad/Education/Undergraduate%20Education/(Link2)%20Core%20curriculum.pdf).
- This handbook addresses these learning outcomes and aims to equip you with the knowledge and skills to practise competently and safely as a junior doctor.

Essential Clinical Skills

 Detailed history taking and examination provide important diagnostic clues in the assessment of skin problems.

Learning outcomes:

- 1. Ability to take a dermatological history
- 2. Ability to explore a patient's concerns and expectations
- 3. Ability to interact sensitively with people with skin disease
- 4. Ability to examine skin, hair, nails and mucous membranes systematically showing respect for the patient
- 5. Ability to describe physical signs in skin, hair, nails and mucosa
- 6. Ability to record findings accurately in patient's records

Taking a dermatological history

- Using the standard structure of history taking, below are the important points to consider when taking a history from a patient with a skin problem (Table 1).
- For dark lesions or moles, pay attention to questions marked with an asterisk (*).

Table 1. Taking a dermatological history

Main headings Key questions	
Presenting complaint	Nature, site and duration of problem
History of presenting complaint	Initial appearance and evolution of lesion*
	Symptoms (particularly itch and pain)*
	Aggravating and relieving factors
	Previous and current treatments (effective or not)
	Recent contact, stressful events, illness and travel
	History of sunburn and use of tanning machines*
	Skin type (see page 70)*
Past medical history	History of atopy i.e. asthma, allergic rhinitis, eczema
	History of skin cancer and suspicious skin lesions
Family history	Family history of skin disease*
Social history	Occupation (including skin contacts at work)
	Improvement of lesions when away from work
Medication and allergies	Regular, recent and over-the-counter medications
Impact on quality of life	Impact of skin condition and concerns

Examining the skin

There are four important principles in performing a good examination of the skin:
 INSPECT, DESCRIBE, PALPATE and SYSTEMATIC CHECK (Table 2).

Table 2. Examining the skin

Main principles	Key features
INSPECT in general	General observation
	Site and number of lesion(s)
	If multiple, pattern of distribution and configuration
DESCRIBE the individual lesion	<u>SCAM</u>
	$\underline{\mathbf{S}}$ ize (the widest diameter), $\underline{\mathbf{S}}$ hape
	<u>C</u> olour
	<u>A</u> ssociated secondary change
	<u>M</u> orphology, <u>M</u> argin (border)

^{*}If the lesion is pigmented, remember **ABCD**

(the presence of any of these features increase the likelihood of melanoma):

Asymmetry (lack of mirror image in any of the four quadrants)

Irregular Border

Two or more Colours within the lesion

Diameter > 6mm

PALPATE the individual lesion

Surface

Consistency

Mobility

SYSTEMATIC CHECK Examine the nails, scalp, hair & mucous membranes

General examination of all systems

Tenderness

Temperature

Communicating examination findings

• In order to describe, record and communicate examination findings accurately, it is important to learn the appropriate terminology (Tables 3-10).

Table 3. General terms

Terms	Meaning	
Pruritus	Itching	
Lesion	An area of altered skin	
Rash	An eruption	
Naevus	A localised malformation of tissue structures	
	Example: (Picture Source: D@nderm)	



Pigmented melanocytic naevus (mole)

Comedone

A plug in a sebaceous follicle containing altered sebum, bacteria and cellular debris; can present as either open (blackheads) or closed (whiteheads)

Example:





Open comedones (left) and closed comedones (right) in acne

Table 4. Distribution (the pattern of spread of lesions)

Terms	Meaning	
Generalised	All over the body	
Widespread	Extensive	
Localised	Restricted to one area of skin only	
Flexural	Body folds i.e. groin, neck, behind ears, popliteal and antecubital fossa	
Extensor	Knees, elbows, shins	
Pressure areas	Sacrum, buttocks, ankles, heels	
Dermatome	An area of skin supplied by a single spinal nerve	
Photosensitive	Affects sun-exposed areas such as face, neck and back of hands	

Example:



Sunburn

Köebner A linear eruption arising at site of trauma

phenomenon Example:



Psoriasis

Table 5. Configuration (the pattern or shape of grouped lesions)

Terms	Meaning	
Discrete	Individual lesions separated from each other	
Confluent	Lesions merging together	
Linear	In a line	
Target	Concentric rings (like a dartboard)	
	Example:	



Erythema multiforme

Annular

Like a circle or ring

Example:



Tinea corporis ('ringworm')

Discoid / A coin-shaped/round lesion

Nummular Example:



Discoid eczema

Table 6. Colour

Terms Meaning

Erythema

Redness (due to inflammation and vasodilatation) which blanches on pressure

Example:



Palmar erythema

Purpura

Red or purple colour (due to bleeding into the skin or mucous membrane) which does not blanch on pressure – petechiae (small pinpoint macules) and ecchymoses (larger bruise-like patches)

Example:



Henoch-Schönlein purpura (palpable small vessel vasculitis)

Hypo- Area(s) of paler skin

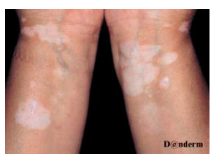
pigmentation Example:



Pityriasis versicolor (a superficial fungus infection)

De- White skin due to absence of melanin

pigmentation Example:



Vitiligo (loss of skin melanocytes)

Hyper- Darker skin which may be due to various causes (e.g. post-inflammatory)

pigmentation Example:



Melasma (increased melanin pigmentation)

Table 7. Morphology (the structure of a lesion) – Primary lesions

Terms Meaning

Macule

A flat area of altered colour

Example:



Freckles

Patch Larger flat area of altered colour or texture Example:



Vascular malformation (naevus flammeus / 'port wine stain')

Papule Solid raised lesion < 0.5cm in diameter Example:



Xanthomata

Nodule Solid raised lesion >0.5cm in diameter with a deeper component

Example: (Picture source: D@nderm)



Pyogenic granuloma (granuloma telangiectaticum)

Plaque Palpable scaling raised lesion >0.5cm in diameter Example:



Psoriasis

Vesicle Raised, clear fluid-filled lesion <0.5cm in diameter (small blister) Example:



Acute hand eczema (pompholyx)

Bulla Raised, clear fluid-filled lesion >0.5cm in diameter
(large blister) Example:



Reaction to insect bites

Pustule Pus-containing lesion < 0.5cm in diameter

Example:



Acne

AbscessLocalised accumulation of pus in the dermis or subcutaneous tissues

Example:



Periungual abscess (acute paronychia)

W(h)eal Transient raised lesion due to dermal oedema Example:



Urticaria

Boil/Furuncle Staphylococcal infection around or within a hair follicle

Carbuncle Staphylococcal infection of adjacent hair follicles (multiple boils/furuncles)

Table 8. Morphology - Secondary lesions (lesions that evolve from primary lesions)

Terms	Meaning
Excoriation	Loss of epidermis following trauma
	Example:



Excoriations in eczema

Lichenification Well-defined roughening of skin with accentuation of skin markings Example:



Lichenification due to chronic rubbing in eczema

Scales Flakes of stratum corneum Example:



Psoriasis (showing silvery scales)

Crust

Rough surface consisting of dried serum, blood, bacteria and cellular debris that has exuded through an eroded epidermis (e.g. from a burst blister) Example:



Impetigo

Scar

New fibrous tissue which occurs post-wound healing, and may be atrophic (thinning), hypertrophic (hyperproliferation within wound boundary), or keloidal (hyperproliferation beyond wound boundary)

Example:



Keloid scars

Ulcer

Loss of epidermis and dermis (heals with scarring)





Leg ulcers

Fissure An epidermal crack often due to excess dryness Example:



Eczema

Striae

Linear areas which progress from purple to pink to white, with the histopathological appearance of a scar (associated with excessive steroid usage and glucocorticoid production, growth spurts and pregnancy) Example:



Striae

Table 9. Hair

Terms Meaning

Alopecia

Loss of hair

Example:



Alopecia areata (well-defined patch of complete hair loss)

Hirsutism Androgen-dependent hair growth in a female

Example:



Hirsutism

Hypertrichosis Non-androgen dependent pattern of excessive hair growth (e.g. in pigmented naevi)

Example:



Hypertrichosis

Table 10. Nails

Terms Meaning

Clubbing

Loss of angle between the posterior nail fold and nail plate
(associations include suppurative lung disease, cyanotic heart disease,
inflammatory bowel disease and idiopathic)

Example: (Picture source: D@nderm)



Clubbing

Koilonychia

Spoon-shaped depression of the nail plate

(associations include iron-deficiency anaemia, congenital and idiopathic)

Example: (Picture source: D@nderm)



Koilonychia

Onycholysis

Separation of the distal end of the nail plate from nail bed (associations include trauma, psoriasis, fungal nail infection and hyperthyroidism)

Example: (Picture source: D@nderm)



Onycholysis

Pitting

Punctate depressions of the nail plate

(associations include psoriasis, eczema and alopecia areata)

Example: (Picture source: D@nderm)



Pitting

Background Knowledge

 This section covers the basic knowledge of normal skin structure and function required to help understand how skin diseases occur.

Learning outcomes:

- 1. Ability to describe the functions of normal skin
- 2. Ability to describe the structure of normal skin
- 3. Ability to describe the principles of wound healing
- 4. Ability to describe the difficulties, physical and psychological, that may be experienced by people with chronic skin disease

Functions of normal skin

- These include:
 - i) Protective barrier against environmental insults
 - ii) Temperature regulation
 - iii) Sensation
 - iv) Vitamin D synthesis
 - v) Immunosurveillance
 - vi) Appearance/cosmesis

Structure of normal skin and the skin appendages

 The skin is the largest organ in the human body. It is composed of the epidermis and dermis overlying subcutaneous tissue. The skin appendages (structures formed by skin-derived cells) are hair, nails, sebaceous glands and sweat glands.

Epidermis

The epidermis is composed of 4 major cell types, each with specific functions (Table
 11).

Table 11. Main functions of each cell type in the epidermis

Cell types	Main functions
Keratinocytes	Produce keratin as a protective barrier
Langerhans' cells	Present antigens and activate T-lymphocytes for immune protection
Melanocytes	Produce melanin, which gives pigment to the skin and protects the
	cell nuclei from ultraviolet (UV) radiation-induced DNA damage
Merkel cells	Contain specialised nerve endings for sensation

 There are 4 layers in the epidermis (Table 12), each representing a different stage of maturation of the keratinocytes. The average epidermal turnover time (migration of cells from the basal cell layer to the horny layer) is about 30 days.

Table 12. Composition of each epidermal layer

Epidermal layers	Composition
Stratum basale	Actively dividing cells, deepest layer
(Basal cell layer)	
Stratum spinosum	Differentiating cells
(Prickle cell layer)	
Stratum granulosum	So-called because cells lose their nuclei and contain
(Granular cell layer)	granules of keratohyaline. They secrete lipid into the
	intercellular spaces.
Stratum corneum	Layer of keratin, most superficial layer
(Horny layer)	

- In areas of thick skin such as the sole, there is a fifth layer, stratum lucidum, beneath the stratum corneum. This consists of paler, compact keratin.
- Pathology of the epidermis may involve:
 - a) changes in epidermal turnover time e.g. psoriasis (reduced epidermal turnover time)
 - b) changes in the surface of the skin or loss of epidermis e.g. scales, crusting, exudate, ulcer
 - c) changes in pigmentation of the skin e.g. hypo- or hyper-pigmented skin

Dermis

- The dermis is made up of collagen (mainly), elastin and glycosaminoglycans, which
 are synthesised by fibroblasts. Collectively, they provide the dermis with strength
 and elasticity.
- The dermis also contains immune cells, nerves, skin appendages as well as lymphatic and blood vessels.
- Pathology of the dermis may involve:
 - a) changes in the contour of the skin or loss of dermis e.g. formation of papules, nodules, skin atrophy and ulcers
 - b) disorders of skin appendages e.g. disorders of hair, acne (disorder of sebaceous glands)
 - c) changes related to lymphatic and blood vessels e.g. erythema (vasodilatation), urticaria (increased permeability of capillaries and small venules), purpura (capillary leakage)

Hair

- There are 3 main types of hair:
 - a) lanugo hair (fine long hair in fetus)
 - b) vellus hair (fine short hair on all body surfaces)
 - c) terminal hair (coarse long hair on the scalp, eyebrows, eyelashes and pubic areas)
- Each hair consists of modified keratin and is divided into the hair shaft (a keratinized tube) and hair bulb (actively dividing cells, and melanocytes which give pigment to the hair).
- Each hair follicle enters its own growth cycle. This occurs in 3 main phases:
 - a) anagen (long growing phase)
 - b) catagen (short regressing phase)
 - c) telogen (resting/shedding phase)
- Pathology of the hair may involve:
 - a) reduced or absent melanin pigment production e.g. grey or white hair
 - b) changes in duration of the growth cycle e.g. hair loss (premature entry of hair follicles into the telogen phase)
 - c) shaft abnormalities

Nails

- The nail is made up of a nail plate (hard keratin) which arises from the nail matrix at the posterior nail fold, and rests on the nail bed.
- The nail bed contains blood capillaries which gives the pink colour of the nails.
- Pathology of the nail may involve:
 - a) abnormalities of the nail matrix e.g. pits and ridges
 - b) abnormalities of the nail bed e.g. splinter haemorrhage
 - c) abnormalities of the nail plate e.g. discoloured nails, thickening of nails

Sebaceous glands

- Sebaceous glands produce sebum via hair follicles (collectively called a
 pilosebaceous unit). They secrete sebum onto the skin surface which lubricates and
 waterproofs the skin.
- Sebaceous glands are stimulated by the conversion of androgens to dihydrotestosterone and therefore become active at puberty.
- Pathology of sebaceous glands may involve:
 - a) increased sebum production and bacterial colonisation e.g. acne
 - b) sebaceous gland hyperplasia

Sweat glands

- Sweat glands regulate body temperature and are innervated by the sympathetic nervous system.
- They are divided into two types: eccrine and apocrine sweat glands.
- Eccrine sweat glands are universally distributed in the skin.
- Apocrine sweat glands are found in the axillae, areolae, genitalia and anus, and modified glands are found in the external auditory canal. They only function from puberty onwards and action of bacteria on the sweat produces body odour.
- Pathology of sweat glands may involve:
 - a) inflammation/infection of apocrine glands e.g. hidradenitis suppurativa
 - b) overactivity of eccrine glands e.g. hyperhidrosis

Principles of wound healing

 Wound healing occurs in 4 phases: haemostasis, inflammation, proliferation and remodelling (Table 13).

Table 13. Stages of wound healing

Stages of wound healing	Mechanisms
Haemostasis	 Vasoconstriction and platelet aggregation
	• Clot formation
Inflammation	Vasodilatation
	 Migration of neutrophils and macrophages
	 Phagocytosis of cellular debris and invading
	bacteria
Proliferation	 Granulation tissue formation (synthesised by
	fibroblasts) and angiogenesis
	 Re-epithelialisation (epidermal cell proliferation
	and migration)
Remodelling	 Collagen fibre re-organisation
	Scar maturation
	• Scar maturation

Emergency Dermatology

- These are rapidly progressive skin conditions and some are potentially lifethreatening. Early recognition is important to implement prompt supportive care and therapy.
- Some are drug reactions and the offending drug should be withdrawn.
- The essential management for all dermatological emergencies, like any emergency, consists of:
 - i) full supportive care ABC of resuscitation
 - ii) withdrawal of precipitating agents
 - iii) management of associated complications
 - iv) specific treatment (highlighted below under each condition)

Learning outcomes:

- 1. Ability to recognise and describe these skin reactions:
 - urticaria
 - erythema nodosum
 - erythema multiforme
- 2. Ability to recognise these emergency presentations, discuss the causes, potential complications and provide first contact care in these emergencies:
 - anaphylaxis and angioedema
 - toxic epidermal necrolysis
 - Stevens-Johnson syndrome
 - acute meningococcaemia
 - erythroderma
 - eczema herpeticum
 - necrotising fasciitis

Urticaria, Angioedema and Anaphylaxis

Causes

 Idiopathic, food (e.g. nuts, sesame seeds, shellfish, dairy products), drugs (e.g. penicillin, contrast media, non-steroidal antiinflammatory drugs (NSAIDs), morphine, angiotensin-converting enzyme inhibitors (ACE-i)), insect bites, contact (e.g. latex), viral or parasitic infections, autoimmune, and hereditary (in some cases of angioedema)

Description

Urticaria is due to a local increase in permeability of capillaries
and small venules. A large number of inflammatory mediators
(including prostaglandins, leukotrienes, and chemotactic factors)
play a role but histamine derived from skin mast cells appears to
be the major mediator. Local mediator release from mast cells can
be induced by immunological or non-immunological mechanisms.

Presentation

- Urticaria (swelling involving the superficial dermis, raising the epidermis): itchy wheals
- Angioedema (deeper swelling involving the dermis and subcutaneous tissues): swelling of tongue and lips
- Anaphylaxis (also known as anaphylactic shock): bronchospasm, facial and laryngeal oedema, hypotension; can present initially with urticaria and angioedema

Management

- Antihistamines for urticaria
- Corticosteroids for severe acute urticaria and angioedema
- Adrenaline, corticosteroids and antihistamines for anaphylaxis

Complications

- Urticaria is normally uncomplicated
- Angioedema and anaphylaxis can lead to asphyxia, cardiac arrest and death



Urticaria



Angioedema

Erythema nodosum

Description

• A hypersensitivity response to a variety of stimuli

Causes

 Group A beta-haemolytic streptococcus, primary tuberculosis, pregnancy, malignancy, sarcoidosis, inflammatory bowel disease (IBD), chlamydia and leprosy

Presentation

- Discrete tender nodules which may become confluent
- Lesions continue to appear for 1-2 weeks and leave bruise-like discolouration as they resolve
- Lesions do not ulcerate and resolve without atrophy or scarring
- The shins are the most common site



Erythema nodosum

Erythema multiforme, Stevens-Johnson syndrome and Toxic epidermal necrolysis

Description

- Erythema multiforme, often of unknown cause, is an acute selflimiting inflammatory condition with herpes simplex virus being the main precipitating factor. Other infections and drugs are also causes. Mucosal involvement is absent or limited to only one mucosal surface.
- Stevens-Johnson syndrome is characterised by mucocutaneous necrosis with at least two mucosal sites involved. Skin involvement may be limited or extensive. Drugs or combinations of infections or drugs are the main associations. Epithelial necrosis with few inflammatory cells is seen on histopathology. The extensive necrosis distinguishes Stevens-Johnson syndrome from erythema multiforme. Stevens-Johnson syndrome may have features overlapping with toxic epidermal necrolysis including a prodromal illness.
- Toxic epidermal necrosis which is usually drug-induced, is
 an acute severe similar disease characterised by extensive skin and
 mucosal necrosis accompanied by systemic toxicity. On
 histopathology there is full thickness epidermal necrosis with
 subepidermal detachment.

Management

- Early recognition and call for help
- Full supportive care to maintain haemodynamic equilibrium

Complications

 Mortality rates are 5-12% with SJS and >30% with TEN with death often due to sepsis, electrolyte imbalance or multi-system organ failure



Erythema multiforme



Stevens-Johnson syndrome

Acute meningococcaemia

Presentation

Description ● A serious communicable infection transmitted via respiratory

secretions; bacteria get into the circulating blood

Cause ● Gram negative diplococcus Neisseria *meningitides*

• Features of meningitis (e.g. headache, fever, neck stiffness),

septicaemia (e.g. hypotension, fever, myalgia) and a typical rash

• Non-blanching purpuric rash on the trunk and extremities, which

may be preceded by a blanching maculopapular rash, and can

rapidly progress to ecchymoses, haemorrhagic bullae and tissue

necrosis

Management ● Antibiotics (e.g. benzylpenicillin)

• Prophylactic antibiotics (e.g. rifampicin) for close contacts (ideally

within 14 days of exposure)

Complications • Septicaemic shock, disseminated intravascular coagulation, multi-

organ failure and death

Erythroderma ('red skin')

Description

• Exfoliative dermatitis involving at least 90% of the skin surface

Causes

 Previous skin disease (e.g. eczema, psoriasis), lymphoma, drugs (e.g.sulphonamides, gold, sulphonylureas, penicillin, allopurinol, captopril) and idiopathic

Presentation

- Skin appears inflamed, oedematous and scaly
- Systemically unwell with lymphadenopathy and malaise

Management

- Treat the underlying cause, where known
- Emollients and wet-wraps to maintain skin moisture
- Topical steroids may help to relieve inflammation

Complications

 Secondary infection, fluid loss and electrolyte imbalance, hypothermia, high-output cardiac failure and capillary leak syndrome (most severe)

Prognosis

- Largely depends on the underlying cause
- Overall mortality rate ranges from 20 to 40%



Erythroderma

Eczema herpeticum (Kaposi's varicelliform eruption)

• Widespread eruption - serious complication of atopic eczema or

less commonly other skin conditions

Cause ● Herpes simplex virus

PresentationExtensive crusted papules, blisters and erosions

• Systemically unwell with fever and malaise

Management ● Antivirals (e.g. aciclovir)

• Antibiotics for bacterial secondary infection

Complications • Herpes hepatitis, encephalitis, disseminated intravascular

coagulation (DIC) and rarely, death



Eczema herpeticum

Necrotising fasciitis

Description ● A rapidly spreading infection of the deep fascia with secondary

tissue necrosis

• Group A haemolytic streptococcus, or a mixture of anaerobic and

aerobic bacteria

• Risk factors include abdominal surgery and medical co-morbidities

(e.g. diabetes, malignancy)

• 50% of cases occur in previously healthy individuals

Presentation ● Severe pain

• Erythematous, blistering, and necrotic skin

• Systemically unwell with fever and tachycardia

• Presence of crepitus (subcutaneous emphysema)

• X-ray may show soft tissue gas (absence should not exclude the

diagnosis)

Management ■ Urgent referral for extensive surgical debridement

• Intravenous antibiotics

Further reading: Hasham S, Matteucci P, Stanley PRW, Hart NB. Necrotising fasciitis. BMJ 2005;330:830-833 (http://www.bmj.com/cgi/content/full/330/7495/830)

Skin Infections / Infestations

- The normal skin microflora and antimicrobial peptides protect the skin against infection. However, when there is skin damage, microorganisms can penetrate resulting in infection.
- There are 3 main types of skin infections according to their sources: bacterial (e.g. staphylococcal and streptococcal), viral (e.g. human papilloma virus, herpes simplex (see page 34) and herpes zoster (see below)), and fungal (e.g. tinea (see page 39 & 40), candida (see page 39 & 40) and yeasts). Infestations (e.g. scabies (see page 58 & 59), cutaneous leishmaniasis) can also occur.



Herpes zoster (shingles) infection due to varicella-zoster virus affecting the distribution of the ophthalmic division of the fifth cranial (trigeminal) nerve Note: Examination for eye involvement is important

Learning outcomes:

Ability to describe the presentation, investigation and management of:

- cellulitis and erysipelas
- staphylococcal scalded skin syndrome
- superficial fungal infections

Erysipelas and Cellulitis

Description

- Spreading bacterial infection of the skin
- Cellulitis involves the deep subcutaneous tissue
- Erysipelas is an acute superficial form of cellulitis and involves the dermis and upper subcutaneous tissue

Causes

- Streptococcus pyogenes and Staphylococcus aureus
- Risk factors include immunosuppression, wounds, leg ulcers, toeweb intertrigo, and minor skin injury

Presentation

- Most common in the lower limbs
- Local signs of inflammation swelling (tumor), erythema (rubor),
 warmth (calor), pain (dolor); may be associated with lymphangitis
- Systemically unwell with fever, malaise or rigors, particularly with erysipelas
- Erysipelas is distinguished from cellulitis by a well-defined, red raised border

Management

- Antibiotics (e.g. flucloxacillin or benzylpenicillin)
- Supportive care including rest, leg elevation, sterile dressings and analgesia

Complications

• Local necrosis, abscess and septicaemia



Cellulitis with elephantiasis of the penis



Erysipelas

Staphylococcal scalded skin syndrome

Description

• Commonly seen in infancy and early childhood

Cause

Production of a circulating epidermolytic toxin from phage group
 II, benzylpenicillin-resistant (coagulase positive) staphylococci

Presentation

- Develops within a few hours to a few days, and may be worse over the face, neck, axillae or groins
- A scald-like skin appearance is followed by large flaccid bulla
- Perioral crusting is typical
- There is intraepidermal blistering in this condition
- Lesions are very painful
- Sometimes the eruption is more localised
- Recovery is usually within 5-7 days

Management

- Antibiotics (e.g. a systemic penicillinase-resistant penicillin, fusidic acid, erythromycin or appropriate cephalosporin)
- Analgesia





Staphylococcal scalded skin syndrome

Superficial fungal infections

Description

 A common and mild infection of the superficial layers of the skin, nails and hair, but can be severe in immunocompromised individuals

Cause

 Three main groups: dermatophytes (tinea/ringworm), yeasts (e.g. candidiasis, malassezia), moulds (e.g. aspergillus)

Presentation

- Varies with the site of infection; usually unilateral and itchy
- Tinea corporis (tinea infection of the trunk and limbs) Itchy, circular or annular lesions with a clearly defined, raised and scaly edge is typical
- Tinea cruris (tinea infection of the groin and natal cleft) very itchy, similar to tinea corporis
- Tinea pedis (athlete's foot) moist scaling and fissuring in toewebs, spreading to the sole and dorsal aspect of the foot
- Tinea manuum (tinea infection of the hand) scaling and dryness in the palmar creases
- Tinea capitis (scalp ringworm) patches of broken hair, scaling and inflammation
- Tinea unguium (tinea infection of the nail) yellow discolouration, thickened and crumbly nail
- Tinea incognito (inappropriate treatment of tinea infection with topical or systemic corticosteroids) – III-defined and less scaly lesions
- Candidiasis (candidal skin infection) white plaques on mucosal areas, erythema with satellite lesions in flexures
- Pityriasis/Tinea versicolor (infection with Malassezia furfur) scaly pale brown patches on upper trunk that fail to tan on sun exposure, usually asymptomatic

Management

- Establish the correct diagnosis by skin scrapings, hair or nail clippings (for dermatophytes); skin swabs (for yeasts)
- General measures: treat known precipitating factors (e.g. underlying immunosuppressive condition, moist environment)

- Topical antifungal agents (e.g. terbinafine cream)
- Oral antifungal agents (e.g. itraconazole) for severe, widespread, or nail infections
- Avoid the use of topical steroids can lead to tinea incognito
- Correct predisposing factors where possible (e.g. moist environment, underlying immunosuppression)



Tinea corporis



Tinea capitis



Tinea manuum (right hand)



Tinea pedis with associated tinea unguium



Candidiasis (right axilla)



Pityriasis versicolor

Skin Cancer

- Skin cancer is one of the most common cancers.
- In general, skin cancer can be divided into: non-melanoma (basal cell carcinoma and squamous cell carcinoma) and melanoma (malignant melanoma).
- Malignant melanoma is the most life-threatening type of skin cancer and is one of the few cancers affecting the younger population.
- Sun exposure is the single most preventable risk factor for skin cancer.

Learning outcomes:

Ability to recognise:

- basal cell carcinoma
- squamous cell carcinoma
- malignant melanoma

Basal cell carcinoma

Description

- A slow-growing, locally invasive malignant tumour of the epidermal keratinocytes normally in older individuals, only rarely metastasises
- Most common malignant skin tumour

Causes

 Risk factors include UV exposure, history of frequent or severe sunburn in childhood, skin type I (always burns, never tans), increasing age, male sex, immunosuppression, previous history of skin cancer, and genetic predisposition

Presentation

- Various morphological types including nodular (most common), superficial (plaque-like), cystic, morphoeic (sclerosing), keratotic and pigmented
- Nodular basal cell carcinoma is a small, skin-coloured papule or nodule with surface telangiectasia, and a pearly rolled edge; the lesion may have a necrotic or ulcerated centre (rodent ulcer)
- Most common over the head and neck

Management

- Surgical excision treatment of choice as it allows histological examination of the tumour and margins
- Mohs micrographic surgery (i.e. excision of the lesion and tissue borders are progressively excised until specimens are microscopically free of tumour) - for high risk, recurrent tumours
- Radiotherapy when surgery is not appropriate
- Other e.g. cryotherapy, curettage and cautery, topical photodynamic therapy, and topical treatment (e.g. imiquimod cream) - for small and low-risk lesions

Complications

Prognosis

- Local tissue invasion and destruction
- Depends on tumour size, site, type, growth pattern/histological subtype, failure of previous treatment/recurrence, and immunosuppression



Basal cell carcinoma - nodular type

Squamous cell carcinoma

Description

 A locally invasive malignant tumour of the epidermal keratinocytes or its appendages, which has the potential to metastasise

Causes

 Risk factors include excessive UV exposure, pre-malignant skin conditions (e.g. actinic keratoses), chronic inflammation (e.g. leg ulcers, wound scars), immunosuppression and genetic predisposition

Presentation

• Keratotic (e.g. scaly, crusty), ill-defined nodule which may ulcerate

Management

• Surgical excision - treatment of choice

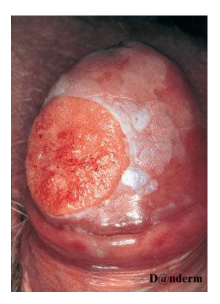
 Mohs micrographic surgery – may be necessary for ill-defined, large, recurrent tumours

• Radiotherapy - for large, non-resectable tumours

Prognosis

 Depends on tumour size, site, histological pattern, depth of invasion, perineural involvement, and immunosuppression





Squamous cell carcinoma – adjacent to ear (left) and glans penis (right)

Malignant melanoma

Description

An invasive malignant tumour of the epidermal melanocytes,
 which has the potential to metastasise

Causes

 Risk factors include excessive UV exposure, skin type I (always burns, never tans), history of multiple moles or atypical moles, and family history or previous history of melanoma

Presentation

• The 'ABCDE Symptoms' rule (*major suspicious features):

Asymmetrical shape*

Border irregularity

Colour irregularity*

Diameter > 6mm

Evolution of lesion (e.g. change in size and/or shape)*

Symptoms (e.g. bleeding, itching)

Types

- More common on the legs in women and trunk in men
- Superficial spreading melanoma common on the lower limbs, in young and middle-aged adults; related to intermittent highintensity UV exposure
- Nodular melanoma common on the trunk, in young and middleaged adults; related to intermittent high-intensity UV exposure
- Lentigo maligna melanoma common on the face, in elderly population; related to long-term cumulative UV exposure
- Acral lentiginous melanoma common on the palms, soles and nail beds, in elderly population; no clear relation with UV exposure

Management

- Surgical excision definitive treatment
- Radiotherapy may sometimes be useful
- Chemotherapy for metastatic disease

Prognosis

- Recurrence of melanoma based on Breslow thickness (thickness of tumour): <0.76mm thick – low risk, 0.76mm-1.5mm thick – medium risk, >1.5mm thick – high risk
- 5-year survival rates based on the TNM classification (primary Tumour, regional Nodes, Metastases): stage 1 (T <2mm thick, N0, M0) 90%, stage 2 (T>2mm thick, N0, M0) 80%, stage 3 (N≥1, M0) 40-50%, and stage 4 (M≥1) 20-30%



Superficial spreading melanoma



Nodular melanoma



Lentigo maligna melanoma



Acral lentiginous melanoma

Inflammatory Skin Conditions

- Eczema, acne and psoriasis are chronic inflammatory skin disorders that follow a relapsing and remitting course. There are many types of eczema but we shall just consider atopic eczema here.
- These skin disorders are not infectious.
- Management is aimed at achieving control and not providing a cure.
- Complications are mainly due to the psychological and social effects.
- Patient education is important in these chronic skin conditions and should concentrate on providing information about the nature of condition, aims of treatment and the available treatment options.

Learning outcomes:

Ability to describe the presentation, demonstrate assessment, formulate a differential diagnosis, instigate investigation and discuss how to provide continuing care of:

- atopic eczema
- acne
- psoriasis

Atopic eczema

Description

- Eczema (or dermatitis) is characterized by papules and vesicles on an erythematous base
- Atopic eczema is the most common type usually develops by early childhood and resolves during teenage years (but may recur)

Epidemiology

Causes

- 20% prevalence in <12 years old in the UK
- Not fully understood, but a positive family history of atopy (i.e. eczema, asthma, allergic rhinitis) is often present
- A primary genetic defect in skin barrier function (loss of function variants of the protein filaggrin) appears to underlie atopic eczema
- Exacerbating factors such as infections, allergens (e.g. chemicals, food, dust, pet fur), sweating, heat and severe stress

Presentation

- Commonly present as itchy, erythematous dry scaly patches
- More common on the face and extensor aspects of limbs in infants, and the flexor aspects in children and adults
- Acute lesions are erythematous, vesicular and weepy (exudative)
- Chronic scratching/rubbing can lead to excoriations and lichenification
- May show nail pitting and ridging of the nails

Management

- General measures avoid known exacerbating agents, frequent emollients +/- bandages and bath oil/soap substitute
- Topical therapies topical steroids for flare-ups; topical immunomodulators (e.g. tacrolimus, pimecrolimus) can be used as steroid-sparing agents
- Oral therapies antihistamines for symptomatic relief, antibiotics (e.g. flucloxacillin) for secondary bacterial infections, and antivirals (e.g. aciclovir) for secondary herpes infection
- Phototherapy and immunosuppressants (e.g. oral prednisolone, azathioprine, ciclosporin) for severe non- responsive cases

Complications

- Secondary bacterial infection (crusted weepy lesions)
- Secondary viral infection molluscum contagiosum (pearly papules with central umbilication), viral warts and eczema herpeticum (see page 34)





Atopic eczema

Further reading: NICE guidelines. Atopic eczema in children, Dec 2007. http://www.nice.org.uk/Guidance/CG57

Acne vulgaris

Description

• An inflammatory disease of the pilosebaceous follicle

Epidemiology

• Over 80% of teenagers aged 13- 18 years

Causes

- Hormonal (androgen)
- Contributing factors include increased sebum production, abnormal follicular keratinization, bacterial colonization (*Propionibacterium acnes*) and inflammation

Presentation

- Non-inflammatory lesions (mild acne) open and closed comedones (blackheads and whiteheads)
- Inflammatory lesions (moderate and severe acne) papules, pustules, nodules, and cysts
- Commonly affects the face, chest and upper back

Management

- General measures no specific food has been identified to cause acne, treatment needs to be continued for at least 6 weeks to produce effect
- Topical therapies (for mild acne) benzoyl peroxide and topical antibiotics (antimicrobial properties), and topical retinoids (comedolytic and anti-inflammatory properties)
- Oral therapies (for moderate to severe acne) oral antibiotics, and anti-androgens (in females)
- Oral retinoids (for severe acne)

Complications

 Post-inflammatory hyperpigmentation, scarring, deformity, psychological and social effects



Comedones



Papules and nodules

<u>Psoriasis</u>

Description

 A chronic inflammatory skin disease due to hyperproliferation of keratinocytes and inflammatory cell infiltration

Types

- Chronic plaque psoriasis is the most common type
- Other types include guttate (raindrop lesions), seborrhoeic (naso-labial and retro-auricular), flexural (body folds), pustular (palmar-plantar), and erythrodermic (total body redness)

Epidemiology

• Affects about 2% of the population in the UK

Causes

- Complex interaction between genetic, immunological and environmental factors
- Precipitating factors include trauma (which may produce a Köebner phenomenon), infection (e.g. tonsillitis), drugs, stress, and alcohol

Presentation

- Well-demarcated erythematous scaly plaques
- Lesions can sometimes be itchy, burning or painful
- Common on the extensor surfaces of the body and over scalp
- Auspitz sign (scratch and gentle removal of scales cause capillary bleeding)
- 50% have associated nail changes (e.g. pitting, onycholysis)
- 5-8% suffer from associated psoriatic arthropathy symmetrical polyarthritis, asymmetrical oligomonoarthritis, lone distal interphalangeal disease, psoriatic spondylosis, and arthritis mutilans (flexion deformity of distal interphalangeal joints)

Management

- General measures avoid known precipitating factors, emollients to reduce scales
- Topical therapies (for localised and mild psoriasis) vitamin D analogues, topical corticosteroids, coal tar preparations, dithranol, topical retinoids, keratolytics and scalp preparations
- Phototherapy (for extensive disease) phototherapy i.e. UVB and photochemotherapy i.e. psoralen+UVA
- Oral therapies (for extensive and severe psoriasis, or psoriasis with systemic involvement) - methotrexate, retinoids, ciclosporin, mycophenolate mofetil, fumaric acid esters,

and biological agents (e.g. infliximab, etanercept, efalizumab)

Complications

• Erythroderma (see page 33), psychological and social effects



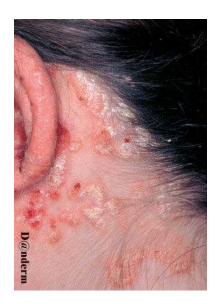




Plaque psoriasis



Nail changes and arthropathy



Scalp involvement

Blistering Disorders

- In general, blistering skin disorders can be divided into: immunobullous diseases
 (e.g. bullous pemphigoid, pemphigus vulgaris), blistering skin infections (e.g. herpes simplex) and other (e.g. porphyria cutanea tarda).
- The fragility of blisters depends on the level of split within the skin an intraepidermal split (a split within the epidermis) causes blisters to rupture easily;
 whereas a sub-epidermal split (a split between the epidermis and dermis) causes
 blisters to be less fragile.
- The common causes of blisters are impetigo (see below), insect bites, herpes simplex infection (see page 34), herpes zoster infection (see page 36), acute contact dermatitis, pompholyx (vesicular eczema of the hands and feet, see below) and burns.
- Bullous pemphigoid (see page 53) and pemphigus vulgaris (see page 54) are uncommon conditions due to immune reaction within the skin.



Bullous impetigo in a new tattoo



Pompholyx

Learning outcomes:

- 1. Ability to recognise common causes of blisters
- 2. Ability to recognise:
 - Bullous pemphigoid
 - Pemphigus vulgaris

Bullous pemphigoid

Description

• A blistering skin disorder which usually affects the elderly

Cause

 Autoantibodies against antigens between the epidermis and dermis causing a sub-epidermal split in the skin

Presentation

- Tense, fluid-filled blisters on an erythematous base
- Lesions are often itchy
- May be preceded by a non-specific itchy rash
- Usually affects the trunk and limbs (mucosal involvement less common)

Management

- General measures wound dressings where required, monitor for signs of infection
- Topical therapies for localised disease topical steroids
- Oral therapies for widespread disease oral steroids, combination
 of oral tetracycline and nicotinamide, immunosuppressive agents
 (e.g. azathioprine, mycophenolate mofetil, methotrexate, and
 other)



Bullous pemphigoid

Pemphigus vulgaris

Description

Cause

- A blistering skin disorder which usually affects the middle-aged
- Autoantibodies against antigens within the epidermis causing an intra-epidermal split in the skin
- **Presentation**
- Flaccid, easily ruptured blisters forming erosions and crusts
- Lesions are often painful
- Usually affects the mucosal areas (can precede skin involvement)
- Management
- General measures wound dressings where required, monitor for signs of infection, good oral care (if oral mucosa is involved)
- Oral therapies high-dose oral steroids, immunosuppressive agents (e.g. methotrexate, azathioprine, cyclophosphamide, mycophenolate mofetil, and other)



Pemphigus vulgaris



Pemphigus vulgaris affecting the oral mucosa

Common Important Problems

- There are several commonly-encountered skin problems in clinical practice. Below are some of the important differential diagnoses for each of these presentations.
- Clinical exposure is the key to achieve competence in diagnosing, investigating and managing these skin problems.

Learning objectives:

Ability to formulate a differential diagnosis, describe the investigation and discuss the management in patients with:

- chronic leg ulcers
- itchy eruption
- a changing pigmented lesion
- purpuric eruption
- a red swollen leg

Chronic leg ulcers

- Leg ulcers are classified according to aetiology. In general, there are three main types: venous, arterial and neuropathic ulcers. Other causes include vasculitic ulcers (purpuric, punched out lesions), infected ulcers (purulent discharge, may have systemic signs) and malignancy (e.g. squamous cell carcinoma in long-standing non-healing ulcers).
- In clinical practice, there can be mixture of arterial, venous and/or neuropathic components in an ulcer.



Neuropathic ulcer



Arterial ulcer



Venous ulcer

Chronic leg ulcers

	Venous ulcer	Arterial IIIcar	Nouronathiculcar
History	- Often painful, worse on standing	- Painful especially at night, worse when	- Often painless
	- History of venous disease e.g. varicose	legs are elevated	- Abnormal sensation
	veins, deep vein thrombosis	- History of arterial disease e.g.	- History of diabetes or neurological disease
		atherosclerosis	
Common sites	- Malleolar area (more common over	- Pressure and trauma sites e.g. pretibial,	- Pressure sites e.g. soles, heel, toes,
	medial than lateral malleolus)	supramalleolar (usually lateral), and at	metatarsal heads
		distal points e.g. toes	
Lesion	- Large, shallow irregular ulcer	- Small, sharply defined deep ulcer	- Variable size and depth
	- Exudative and granulating base	- Necrotic base	- Granulating base
			- May be surrounded by or underneath a
			hyperkeratotic lesion (e.g. callus)
Associated	- Warm skin	- Cold skin	- Warm skin
features	- Normal peripheral pulses	- Weak or absent peripheral pulses	- Normal peripheral pulses*
	- Leg oedema, haemosiderin and melanin	- Shiny pale skin	*cold, weak or absent pulses if it is a
	deposition (brown pigment),	- Loss of hair	neuroischaemic ulcer
	lipodermatosclerosis, and atrophie		- Peripheral neuropathy
	blanche (white scarring with dilated		
	capillaries)		
Possible	- Normal ankle/brachial pressure index	- ABPI < 0.8 - presence of arterial	- ABPI < 0.8 implies a neuroischaemic ulcer
investigations	(i.e. ABPI 0.8-1)	insufficiency	- X-ray to exclude osteomyelitis
		- Doppler studies and angiography	
Management	- Compression bandaging	- Vascular reconstruction	- Wound debridement
	(after excluding arterial insufficiency)	- Compression bandaging is contraindicated	- Regular repositioning, appropriate
			tootwear and good nutrition

Itchy eruption

An itchy (pruritic) eruption can be caused by an inflammatory condition (e.g. eczema), infection (e.g. varicella), infestation (e.g. scabies), allergic reaction (e.g. some cases of urticaria) or an unknown cause, possibly autoimmune (e.g. lichen planus).



tchy eruption

History - Personal or family history of attopy attopy attopy attopy - May have history of contact with symptomatic individual autopy Common sites - Variable (e.g. flexor aspects in allergens, irritants) - Sides of fingers, finger web children and adults with atopic mipples and genitals Lesion - Dry, erythematous patches - Linear burrows (may be carde eczema is erythematous, vesicular and exudative Associated - Secondary bacterial or viral infections - Secondary eczema and impetigo Possible - Patch testing - Skin scrape, extraction of mipples with a proposal and infections Possible - Patch testing - Skin scrape, extraction of mipples and genitals Possible - Serum lgE levels - Skin scrape, extraction of mipples and genitals Possible - Patch testing - Skin scrape, extraction of mipples Romanagement - Emollients - Scrabicide (e.g. permethrin or malathion) - Corticosteroids - Antihistamines	ccema	Olucalia	ricuen pianus
atopy - Exacerbating factors (e.g. allergens, irritants) - Variable (e.g. flexor aspects in children and adults with atopic eczema) - Dry, erythematous patches - Line erythematous, vesicular and exudative - Secondary bacterial or viral infections infections - Patch testing - Secondary bacterial or viral and sand vesions and sexumable - Secondary bacterial or viral infections - Serum IgE levels - Skin swab - Corticosteroids - Corticosteroids - Corticosteroids - Antil - Immunomodulators - Antil	- May have history of contact	- Precipitating factors (e.g. food,	- Family history in 10% of cases
- Exacerbating factors (e.g. allergens, irritants) - Variable (e.g. flexor aspects in children and adults with atopic children and adults with atopic eczema) - Dry, erythematous patches - Linea erythematous, vesicular and exudative - Secondary bacterial or viral infections infections - Patch testing - Serum IgE levels and value - Serum IgE levels - Skin swab - Serum IgE levels - Skin swab - Corticosteroids - Corticosteroids - Antil - Immunomodulators - Antil	with symptomatic individuals	contact, drugs)	- May be drug-induced
allergens, irritants) - Variable (e.g. flexor aspects in children and adults with atopic wrist eczema) - Dry, erythematous patches - Acute eczema is tortue erythematous, vesicular and exudative - Secondary bacterial or viral imperinfections - Patch testing - Patch testing - Serum IgE levels - Skin swab - Serum IgE levels - Shin swab - Corticosteroids - Immunomodulators - Antil			
children and adults with atopic wrist eczema) - Dry, erythematous patches - Lines - Acute eczema is erythematous, vesicular and exudative - Secondary bacterial or viral infections infections - Serum IgE levels - Skin swab - Skin swab - Corticosteroids - Corticosteroids - Corticosteroids - Corticosteroids - Corticosteroids - Corticosteroids - Antil - Antil - Antil - Antil	ts)		
children and adults with atopic wrist eczema) - Dry, erythematous patches - Acute eczema is erythematous, vesicular and exudative - Secondary bacterial or viral imperinfections infections - Patch testing - Patch testing - Serum IgE levels - Skin swab - Serum IgE levels - Skin swab - Corticosteroids - Immunomodulators - Antil	- Sides of fingers, finger webs,	- No specific tendency	- Forearms, wrists, and legs
eczema) nippl - Dry, erythematous patches - Acute eczema is - Acute eczema is erythematous, vesicular and exudative - Secondary bacterial or viral impe infections infections - Patch testing - Serum IgE levels - Skin swab - Serum IgE levels - Skin swab - Corticosteroids - Immunomodulators - Antil	Ilts with atopic wrists, elbows, ankles, feet,		- Always examine the oral
- Dry, erythematous patches - Acute eczema is - Acute eczema is erythematous, vesicular and exudative - Secondary bacterial or viral imperions infections - Patch testing - Patch testing - Skin swab - Skin swab - Skin swab - Corticosteroids - Immunomodulators - Antil	nipples and genitals		mucosa
ed erythematous, vesicular and exudative - Secondary bacterial or viral infections imperiment - Patch testing - Serum IgE levels - Skin swab - Corticosteroids - Immunomodulators - Antil - Immunomodulators - Antil	- Linear burrows (may be	- Pink wheals (transient)	- Violaceous (lilac) flat-topped
ed evythematous, vesicular and exudative - Secondary bacterial or viral imperions infections - Patch testing - Serum IgE levels - Skin swab - Serum IgE levels - Skin swab - Corticosteroids - Corticosteroids - Immunomodulators - Antil - Antil	tortuous) or rubbery nodules	- May be round, annular, or	papules
ed - Secondary bacterial or viral - Seco infections - Secondary bacterial or viral - Seco infections - Secondary bacterial or viral - Secondary bacterial or viral - Secondary - Skin swab - Skin swab - Skin swab - Scab - Corticosteroids - Scab - Immunomodulators - Antil - Antil		polycyclic	- Symmetrical distribution
ed - Secondary bacterial or viral - Seco infections imperations - Patch testing - Skin stions - Serum IgE levels - Skin swab - Skin swab - Skin swab - Corticosteroids - Scab - Corticosteroids - Smullients - Smulli			
infections imperions imperions - Patch testing - Serum IgE levels - Skin swab - Skin swab - Emollients - Corticosteroids - Corticosteroids - Antil - Antil	- Secondary eczema and	- May be associated with	- Nail changes and hair loss
- Patch testing - Serum IgE levels - Skin swab - Emollients - Corticosteroids - Immunomodulators - Antil		angioedema or anaphylaxis	- Lacy white streaks on the oral
- Patch testing - Serum IgE levels - Skin swab - Emollients - Corticosteroids - Immunomodulators - Antil			mucosa and skin lesions
- Patch testing - Serum IgE levels - Skin swab - Emollients - Corticosteroids - Immunomodulators - Antil			(Wickham's striae)
- Serum IgE levels - Skin swab - Emollients - Corticosteroids - Immunomodulators	scrape, extraction of mite	- Bloods and urinalysis to	- Skin biopsy
- Skin swab - Emollients - Corticosteroids - Immunomodulators	and view under microscope	exclude a systemic cause	
- Emollients - Corticosteroids - Immunomodulators			
<u> </u>	1	Antihistamines	- Corticosteroids
	1	Corticosteroids	- Antihistamines
- Antihistamines			

A changing pigmented lesion

A changing pigmented lesion can be benign (e.g. melanocytic naevi, seborrhoeic wart) or malignant (e.g. malignant melanoma).



Malignant melanoma



Congenital naevus

Seborrhoeic keratoses

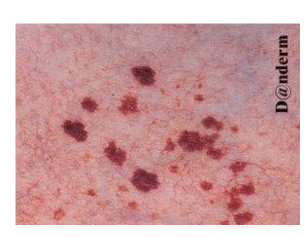


A changing pigmented lesion

History - Not usually preser during infancy, chi - Asymptomatic Common sites - Variable	- Not usually present at birth but develop during infancy, childhood or adolescence - Asymptomatic	Seborrhoeic wart - Tend to arise in the middle-aged or elderly - Often multiple and asymptomatic	Malignant melanoma
1 1 1	a)	Tend to arise in the middle-aged or elderly Often multiple and asymptomatic	
1 1		Often multiple and asymptomatic	- Tend to occur in adults or the middle-aged
			- History of evolution of lesion
			- May be symptomatic (e.g. itchy, bleeding)
			- Presence of risk factors
		- Face and trunk	- More common on the legs in women and
			trunk in men
	- Congenital naevi may be large,	- Warty greasy papules or nodules	- Features of ABCDE :
pigmented, protuberant and hairy		- 'Stuck on' appearance, with well-defined	A symmetrical shape
- Junctional naevi a	- Junctional naevi are small, flat and dark	edges	B order irregularity
- Intradermal naevi	- Intradermal naevi are usually dome-shape		Colour irregularity
papules or nodules	S		D iameter > 6mm
- Compound naevi	- Compound naevi are usually raised, warty,		Evolution of lesion
hyperkeratotic, and/or hairy	d/or hairy		
Management - Rarely needed	1	- Rarely needed	- Excision

Purpuric eruption

- thrombocytopenic purpura) or non-thrombocytopenic e.g. trauma, drugs (e.g. steroids), aged skin, vasculitis (e.g. Henoch-Schönlein purpura). A purpuric eruption can be thrombocytopenic (e.g. meningococcal septicaemia, disseminated intravascular coagulation, idiopathic
- Platelet counts and a clotting screen are important to exclude coagulation disorders.



Senile purpura



Henoch-Schönlein purpura

Purpuric eruption

	Meningococcal septicaemia	Disseminated intravascular	Vasculitis	Senile purpura
		coagulation		
History	- Acute onset	- History of trauma, malignancy,	- Painful lesions	- Arise in the elderly population
	- Symptoms of meningitis and	sepsis, obstetric complications,		with sun-damaged skin
	septicaemia	transfusions, or liver failure		
Common sites	- Extremities	- Spontaneous bleeding from	- Dependent areas (e.g. legs,	- Extensor surfaces of hands
		ear, nose and throat,	buttocks, flanks)	and forearms
		gastrointestinal tract,		- Such skin is easily traumatised
		respiratory tract or wound site		
Lesion	- Petechiae, ecchymoses,	- Petechiae, ecchymoses,	- Palpable purpura (often	- Non-palpable purpura
	haemorrhagic bullae and/or	haemorragic bullae and/or	painful)	- Surrounding skin is atrophic
	tissue necrosis	tissue necrosis		and thin
Associated	- Systemically unwell	- Systemically unwell	- Systemically unwell	- Systemically well
features				
Possible	- Bloods	- Bloods (a clotting screen is	- Bloods and urinalysis	- No investigation is needed
investigations	- Lumbar puncture	important)	- Skin biopsy	
Management	- Antibiotics	- Treat the underlying cause	- Treat the underlying cause	- No treatment is needed
		- Transfuse for coagulation	- Steroids and	
		deficiencies	immunosuppressants if there	
		- Anticoagulants for thrombosis	is systemic involvement	

A red swollen leg

The main differential diagnoses for a red swollen leg are cellulitis, erysipelas, venous thrombosis and chronic venous insufficiency.

	Cellulitis/Erysipelas	Venous thrombosis	Chronic venous insufficiency
History	- Painful spreading rash	- Pain with swelling and redness	- Heaviness or aching of leg, which is
	- History of abrasion or uicer	- History of prolonged bed rest, long haul	worse on standing and relieved by
			- History of venous thrombosis
Lesion	- Erysipelas (well-defined edge)	- Complete venous occlusion may lead to	- Discoloured (blue-purple)
	- Cellulitis (diffuse edge)	cyanotic discolouration	- Oedema (improved in the morning)
			- Venous congestion and varicose veins
Associated	- Systemically unwell with fever and malaise	- Usually systemically well	- Lipodermatosclerosis (erythematous
features	- May have lymphangitis	- May present with pulmonary embolism	induration, creating 'champagne
			bottle' appearance)
			- Stasis dermatitis (eczema with
			inflammatory papules, scaly and
			crusted erosions)
			- Venous ulcer
Possible	- Anti-streptococcal O titre (ASOT)	- D-dimer	- Doppler ultrasound and/or venography
investigations	- Skin swab	- Doppler ultrasound and/or venography	
Management	- Antibiotics	- Anticoagulants	- Leg elevation and compression
			stockings
			- Sclerotherapy or surgery for varicose
			veins

Management

- Treatment modalities for skin disease can be broadly categorised into medical therapy (topical and systemic treatments) and physical therapy (e.g. cryotherapy, phototherapy, photodynamic therapy, lasers and surgery).
- Topical treatments directly deliver treatment to the affected areas and this reduces systemic side effects. It is suitable for localised and less severe skin conditions. They consist of active constituents which are transported into the skin by a base (also known as a 'vehicle'). Examples of active ingredients are steroids, tar, immunomodulators, retinoids, and antibiotics. The common forms of base are lotion (liquid), cream (oil in water), gel (organic polymers in liquid, transparent), ointment (oil with little or no water) and paste (powder in ointment).
- Systemic therapy is used for extensive and more serious skin conditions, if the
 treatment is ineffective topically or if there is systemic involvement. However, they
 have the disadvantage of causing systemic side effects.

Learning objectives:

Ability to describe the principles of use of the following drugs:

- emollients
- topical/oral corticosteroids
- oral aciclovir
- oral antihistamines
- topical/oral antibiotics
- topical antiseptics

Emollients

• Aqueous cream, emulsifying ointment, liquid paraffin and white soft

paraffin in equal parts (50:50)

Quantity ● 500 grams per tub

Indications
 To rehydrate skin and re-establish the surface lipid layer

• Useful for dry, scaling conditions and as soap substitutes

• Reactions may be irritant or allergic (e.g. due to preservatives or perfumes

in creams)

Topical/Oral corticosteroids

Examples ■ Topical steroids: classified as mildly potent (e.g, hydrocortisone),

moderately potent (e.g. clobetasone butyrate (Eumovate)), potent

(e.g. betamethasone valerate (*Betnovate*)), and very potent (e.g. clobetasol

propionate (Dermovate))

• Oral steroids: prednisolone

Quantity • Usually 30 grams per tube (enough to cover the whole body once)

IndicationsAnti-inflammatory and anti-proliferative effects

 \bullet Useful for allergic and immune reactions, inflammatory skin conditions,

blistering disorders, connective tissue diseases, and vasculitis

• Local side effects (from topical corticosteroids): skin atrophy (thinning),

telangiectasia, striae, may mask, cause or exacerbate skin infections,

acne, or perioral dermatitis, and allergic contact dermatitis.

• Systemic side effects (from oral corticosteroids): Cushing's syndrome,

immunosuppression, hypertension, diabetes, osteoporosis, cataract, and

steroid-induced psychosis

Oral aciclovir

Examples • Aciclovir

• Viral infections due to herpes simplex and herpes zoster virus

• Gastrointestinal upsets, raised liver enzymes, reversible neurological

reactions, and haematological disorders

Oral antihistamines

Classified into nonsedative (e.g. cetirizine, loratadine) and sedative

antihistamines (e.g. chlorpheniramine, hydroxyzine)

Indications

- Block histamine receptors producing an anti-pruritic effect
- Useful for type-1 hypersensitivity reactions and eczema (especially sedative antihistamines for children)

Side effects

Sedative antihistamines can cause sedation and anticholinergic effects
 (e.g. dry mouth, blurred vision, urinary retention, and constipation)

Topical/Oral antibiotics

Examples

- Topical antibiotics: fusidic acid, mupirocin (Bactroban), neomycin
- Oral antibiotics: penicillins, cephalosporins, gentamicin, macrolides, nitrofurantoin, quinolones, tetracyclines, vancomycin, metronidazole, trimethoprim

Indications

• Useful for bacterial skin infections, and some are used for acne

Side effects

- Local side effects (from topical antibiotics): local skin irritation/allergy
- Systemic side effects (from oral antibiotics): gastrointestinal upset, rashes, anaphylaxis, vaginal candidiasis, antibiotic-associated infection such as Clostridium difficile, and antibiotic resistance (rapidly appears to fusidic acid)

Topical antiseptics

Examples

• Chlorhexidine, cetrimide, povidone-iodine

Indications

• Treatment and prevention of skin infection

Side effects

• Local side effects: local skin irritation/allergy

Oral retinoids

Examples

• Isotretinoin, Acitretin

Indications

• Acne, psoriasis, and disorders of keratinisation

Side effects

- Mucocutaneous reactions such as dry skin, dry lips and dry eyes,
 disordered liver function, hypercholesterolaemia, hypertriglyceridaemia,
 myalgia, arthralgia and depression
- Teratogenicity: effective contraception must be practised one month before, during and at least one month after isotretinoin, but for two years after Acitretin (consult current BNF for further details)

Practical Skills

- There are four main aspects to focus on in clinical practice:
 - Patient education, particularly on the nature of disease, treatment and ways to achieve full compliance and effectiveness, and prevention strategies
 - ii) Effective written communication to general practitioner so that patients care can be continued appropriately
 - iii) Good prescribing skills
 - iv) Good clinical examination and appropriate investigations to facilitate accurate diagnosis
- This section highlights several general points on the important clinical skills in dermatology.

Learning objectives:

- 1. Ability to perform the following tasks:
 - explain how to use an emollient or a topical corticosteroid
 - make a referral
 - write a discharge letter
 - write a prescription for emollient
 - take a skin swab
 - take a skin scrape
 - measure the ankle-brachial pressure index and interpret the result
- 2. Describe the principles of prevention in:
 - pressure sores
 - sun damage and skin cancer

Patient education

How to use emollients

Apply liberally and regularly

How to use topical corticosteroids

- Apply thinly and only for short-term use (often 1 or 2 weeks only)
- Only use 1% hydrocortisone or equivalent strength on the face
- Fingertip unit (advised on packaging) strip of cream the length of a fingertip

Preventing pressure sores

- Pressure sores are due to ischaemia resulting from localised damage to the skin caused by sustained pressure, friction and moisture, particularly over bony prominences.
- Preventative measures involve frequent repositioning, nutritional support, and use of pressure relieving devices e.g. special beds

Preventing sun damage and skin cancer

- Excessive exposure to UV radiation is the most significant and preventable risk factor for the development of skin cancer (Table 14)
- Skin types I and II are at higher risk of developing skin cancer with excessive sun exposure than other skin types (Table 15)

Table 14. SMART ways to avoid excessive sun exposure

Spend time in the shade between 11am-3pm

Make sure you never burn

Aim to cover up with a t-shirt, wide-brimmed hat and sunglasses

Remember to take extra care with children

Then use Sun Protection Factor (SPF) 30+ sunscreen

Table 15. Skin types

Skin types	Description
1	Always burns, never tans
II	Always burns, sometimes tans
III	Sometimes burns, always tans
IV	Never burns, always tans

Written communication

Writing a referral letter

Important points to include:

- Reason(s) for referral, current presentation, and impact of disease
- Patient's medical and social background
- Current and previous treatment, length of treatment, and response to treatment

Writing a discharge letter

Important points to include:

- Reason(s) for admission and current presentation
- Hospital course
- Investigation results
- Diagnostic impression
- Management plan (including treatment and follow-up appointment)
- Content of patient education given

Prescribing skills

Writing a prescription

General tips:

- Include drug name, dose, frequency and an intended duration/review date
- 30 grams of cream/ointment covers the whole adult body area
- 1 fingertip unit covers the area of two palms and equals ½ gram

Prescribing emollients

General tips

- Emollients come in 500 gram tubs
- In general, ointment-based emollients are useful for dry, scaling skin whereas creams and lotions are for red, inflamed and weeping lesions

Prescribing topical corticosteroids

General tips

- Prescribe the weakest potency corticosteroid that is effective
- Use only for short term
- Need to specify the base i.e. cream, lotion or ointment

Clinical examination and investigations

Taking a skin swab

- Skin swabs can be taken from vesicles, pustules, erosions, ulcers and mucosal surfaces for microbial culture.
- Surface swabs are generally not encouraged.

Taking a skin scrape

• Skin scrapes are taken from scaly lesions by gentle use of a scalpel in suspected fungal infection (to show evidence of fungal hyphae and/or spores) and from burrows in scabies (see page 59).

Measuring ankle-brachial pressure index (ABPI)

- ABPI is used to identify the presence and severity of peripheral arterial insufficiency,
 which is important in the management of leg ulcers.
- Measure the cuff pressure of dorsalis pedis or posterior tibial artery using a Doppler and compare it to the pressure of brachial artery.
- The ABPI is measured by calculating the ratio of highest pressure obtained from the ankle to highest brachial pressure of the two arms, and is normally >0.8.
- Inappropriately high reading will be obtained in calcified vessels (often in diabetics).

Acknowledgements

We wish to acknowledge the following contributors:

- Dr Mark Goodfield, former President (2008-2010) of the British Association of Dermatologists, for writing the Foreword.
- Dr Niels K. Veien for allowing us to use his photographs. All illustrations in this handbook were obtained from "D@nderm" with his permission.
- Dr Susan Burge, retired Consultant Dermatologist, Oxford Radcliffe Hospitals NHS
 Trust, Professor Peter Friedmann, Emeritus Professor of Dermatology, Southampton
 General Hospital, and Professor Lesley Rhodes, Professor of Experimental
 Dermatology, University of Manchester for reviewing and contributing valuable suggestions.
- Mr Kian Tjon Tan, Specialty Registrar in Plastic Surgery, Royal Preston NHS
 Foundation Trust for contributing the chapter Background Knowledge.
- Dr Yi Ning Chiang, Specialty Doctor in Dermatology, Southport and Ormskirk Hospital
 NHS Trust for contributing the chapter Common Important Problems.

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