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SUMMARY
INTRODUCTION

Although common dermatoses are not generally considered to pose a significant threat to the health of an individual, they can become a major problem for an entire armed force. In harsh environments, such as one would encounter on a field of battle, these minor afflictions can become incapacitating, thus rendering an individual unfit for duty. In a large unit, this collective loss of fighting force could seriously impair the effectiveness of the unit as a whole. During the Vietnam conflict, dermatoses were the third-highest cause of hospital admissions for disease, and outpatient dermatology visits were double the number required for any other condition.1,2 Clearly, skin diseases can have a staggering impact on military operations.

### TABLE 20-1

**MOST COMMON DIAGNOSES IN NEW PATIENTS SEEN AT DERMATOLOGY CLINIC, 17TH FIELD HOSPITAL, SAIGON, VIETNAM, JULY 1967 (% of total shown)**

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>No. of Cases (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pyoderma</td>
<td>47 (10.0)</td>
</tr>
<tr>
<td>Miliaria</td>
<td>43 (9.2)</td>
</tr>
<tr>
<td>Tinea</td>
<td>43 (9.2)</td>
</tr>
<tr>
<td>Verrucae</td>
<td>37 (7.9)</td>
</tr>
<tr>
<td>Eczematous dermatitis</td>
<td>26 (5.6)</td>
</tr>
<tr>
<td>Candidiasis</td>
<td>22 (4.7)</td>
</tr>
<tr>
<td>Infected eczematous dermatitis</td>
<td>20 (4.3)</td>
</tr>
<tr>
<td>Acne</td>
<td>18 (3.8)</td>
</tr>
<tr>
<td>Tinea versicolor</td>
<td>15 (3.2)</td>
</tr>
<tr>
<td>Urticaria</td>
<td>13 (2.8)</td>
</tr>
<tr>
<td>Contact dermatitis</td>
<td>11 (2.3)</td>
</tr>
<tr>
<td>Plantar warts</td>
<td>11 (2.3)</td>
</tr>
<tr>
<td>Alopecia areata</td>
<td>10 (2.1)</td>
</tr>
<tr>
<td>Pseudofolliculitis barbae</td>
<td>9 (1.9)</td>
</tr>
<tr>
<td>Psoriasis</td>
<td>7 (1.5)</td>
</tr>
<tr>
<td>Others</td>
<td>137 (29.2)</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>469 (100.0)</strong></td>
</tr>
</tbody>
</table>


### TABLE 20-2

**PROPORTIONATE DISTRIBUTION OF SKIN DISEASES SEEN IN U.S. DERMATOLOGY CLINIC, 95TH EVACUATION HOSPITAL, DANANG, VIETNAM, 15 MAY 1970 TO 31 JULY 1971 (14.5 MONTHS) (% of total shown)**

<table>
<thead>
<tr>
<th>Disease</th>
<th>No. of Cases (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Verrucae, all types</td>
<td>729 (15.83)</td>
</tr>
<tr>
<td>Acne, all types</td>
<td>466 (10.12)</td>
</tr>
<tr>
<td>Dermatophytosis, all types</td>
<td>371 (8.06)</td>
</tr>
<tr>
<td>Pseudofolliculitis barbae</td>
<td>289 (6.28)</td>
</tr>
<tr>
<td>Penile ulcer [? chancroid]</td>
<td>221 (4.80)</td>
</tr>
<tr>
<td>Miliaria</td>
<td>199 (4.32)</td>
</tr>
<tr>
<td>Pyoderma, all types</td>
<td>178 (3.87)</td>
</tr>
<tr>
<td>Contact dermatitis</td>
<td>167 (3.63)</td>
</tr>
<tr>
<td>Urticaria</td>
<td>126 (2.74)</td>
</tr>
<tr>
<td>Tinea versicolor</td>
<td>123 (2.67)</td>
</tr>
<tr>
<td>Psoriasis</td>
<td>106 (2.30)</td>
</tr>
<tr>
<td>Atopic dermatitis</td>
<td>95 (2.06)</td>
</tr>
<tr>
<td>Dyshidrosis</td>
<td>95 (2.06)</td>
</tr>
<tr>
<td>Alopecia areata</td>
<td>82 (1.78)</td>
</tr>
<tr>
<td>Monilia</td>
<td>71 (1.54)</td>
</tr>
<tr>
<td>Lichen planus</td>
<td>70 (1.52)</td>
</tr>
<tr>
<td>Herpes progenitalis</td>
<td>68 (1.48)</td>
</tr>
<tr>
<td>Seborrheic dermatitis</td>
<td>56 (1.22)</td>
</tr>
<tr>
<td>Miscellaneous dermatoses and dermatitides</td>
<td>51 (1.11)</td>
</tr>
<tr>
<td>Insect bites</td>
<td>48 (1.04)</td>
</tr>
<tr>
<td>Molluscum contagiosum</td>
<td>41 (0.89)</td>
</tr>
<tr>
<td>Sebaceous cyst</td>
<td>40 (0.87)</td>
</tr>
<tr>
<td>Pityriasis rosea</td>
<td>39 (0.85)</td>
</tr>
<tr>
<td>Hand and foot eczema</td>
<td>37 (0.80)</td>
</tr>
<tr>
<td>Lichen simplex chronic</td>
<td>35 (0.76)</td>
</tr>
<tr>
<td>Syphilis infection, late and early</td>
<td>33 (0.72)</td>
</tr>
<tr>
<td>Erythema multiforme</td>
<td>32 (0.69)</td>
</tr>
<tr>
<td>Nevi</td>
<td>32 (0.69)</td>
</tr>
<tr>
<td>Balanitis</td>
<td>31 (0.67)</td>
</tr>
<tr>
<td>Basal cell epithelioma</td>
<td>25 (0.54)</td>
</tr>
<tr>
<td>Keloids</td>
<td>24 (0.52)</td>
</tr>
<tr>
<td>Corns and calluses</td>
<td>24 (0.52)</td>
</tr>
<tr>
<td>Drug eruptions</td>
<td>21 (0.45)</td>
</tr>
<tr>
<td>Vitiligo</td>
<td>20 (0.43)</td>
</tr>
<tr>
<td>Photoallergy</td>
<td>15 (0.33)</td>
</tr>
<tr>
<td>Nummular eczema</td>
<td>14 (0.30)</td>
</tr>
<tr>
<td>Pruritus</td>
<td>14 (0.30)</td>
</tr>
<tr>
<td>No diagnosis</td>
<td>56 (1.22)</td>
</tr>
<tr>
<td>Others</td>
<td>461 (10.02)</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>4,605 (100.00)</strong></td>
</tr>
</tbody>
</table>

TABLE 20-3
DERMATOLOGY ADMISSIONS TO A BRITISH GENERAL HOSPITAL DURING WORLD WAR I

<table>
<thead>
<tr>
<th></th>
<th>1915</th>
<th>1916</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Nov</td>
<td>Dec</td>
</tr>
<tr>
<td>Impetigo</td>
<td>122</td>
<td>172</td>
</tr>
<tr>
<td>Scabies</td>
<td>95</td>
<td>770</td>
</tr>
<tr>
<td>Boils</td>
<td>24</td>
<td>59</td>
</tr>
<tr>
<td>Pediculosis</td>
<td>–</td>
<td>17</td>
</tr>
<tr>
<td>Psoriasis</td>
<td>11</td>
<td>24</td>
</tr>
<tr>
<td>Eczema</td>
<td>7</td>
<td>22</td>
</tr>
<tr>
<td>Seborrhoea</td>
<td>8</td>
<td>12</td>
</tr>
<tr>
<td>Acne</td>
<td>6</td>
<td>22</td>
</tr>
<tr>
<td>Dermatitis</td>
<td>7</td>
<td>10</td>
</tr>
<tr>
<td>Syphilis</td>
<td>8</td>
<td>13</td>
</tr>
<tr>
<td>Folliculitis</td>
<td>1</td>
<td>11</td>
</tr>
<tr>
<td>Urticaria</td>
<td>2</td>
<td>4</td>
</tr>
<tr>
<td>Sycosis</td>
<td>4</td>
<td>2</td>
</tr>
<tr>
<td>Pityriasis rosea</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>Erythema</td>
<td>3</td>
<td>7</td>
</tr>
<tr>
<td>Carbuncle</td>
<td>2</td>
<td>4</td>
</tr>
<tr>
<td>Herpes zoster</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Lupus</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Erysipelas</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Ecthyma</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Lichen planus</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Erythema nodosum</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Ichthyosis</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Sudamina</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>


One might expect the largest category of dermatologic casualties to be composed of exotic tropical diseases; however, the greatest number of casualties has been related to common skin maladies. Experience from previous battles clearly indicates that common dermatoses are a major source of ineffectiveness and temporary disability among battlefield personnel. Pillsbury and Livingood\(^3\) reported that the entire group of rare dermatoses was of no significance in the total disability from skin diseases in World War II. Statistics available from Vietnam and World War I (Tables 20-1 through 20-3) support a similar conclusion.\(^3\)\(^4\) It is apparent from these experiences that many casualties can be averted in the future if more military medical personnel are trained in the recognition and management of some common skin diseases.

Tables 20-1 through 20-3 list a number of skin diseases that were discussed in previous chapters, such as impetigo (Chapter 13, Bacterial Skin Diseases), scabies (Chapter 8, Arthropod and Other Animal Bites), and miliaria (Chapter 3, Skin Diseases Associated with Excessive Heat, Humidity, and Sunlight). This chapter will address common dermatoses that did not fit in the previous chapters of this textbook.
Psoriasis is a common, genetically determined, inflammatory skin disease characterized by distinctive lesional morphology and distribution. It affects men and women equally and typically runs a chronic course that is marked by frequent relapses. It occurs worldwide and affects approximately 1% of the population in the United States. Psoriasis can have its onset at any age, but is most likely to appear in early adulthood. Although its exact etiology is unknown, it is clear that psoriatic skin is in a hyperproliferative state with a marked increase in the rate of keratinocyte replication.

Fig. 20-1. Psoriasis. (a) Large, geographic plaques. Note how lesions suggest the coastline of a map. (b) Note symmetry and extensive involvement of lesions. (c) Lichenified geographic plaques in obese individual. (d) Characteristic silvery scale.
Clinical Features

Psoriasis is a classic example of a papulosquamous disease, characterized by erythematous papules and plaques with a silvery scale. The disorder can vary from a focal disease consisting of localized lesions to a widespread eruption and even a generalized erythroderma with exfoliation.\(^\text{11,12}\)

The characteristic early lesion of psoriasis is an erythematous papule with a scale that can be subtle but is usually obvious. Typically, the papules gradually expand and coalesce to form sharply demarcated, symmetrically distributed plaques (Figure 20-1). Frequently, the plaques become surrounded by a thin zone of perilesional blanching, known as a Woronoff ring. This ring is most frequently seen in the early stages of resolution in ultraviolet (UV) light therapy (Figure 20-2). Although some authorities\(^\text{13}\) have shown evidence of prostaglandin inhibition to explain this phenomenon, the precise mechanisms involved are likely to be more complex.\(^\text{14}\)

Psoriasis can occur at any cutaneous site, although it has a predilection for the scalp, elbows, knees, extensor aspects of the extremities, and the nails. It also frequently involves the penis (Figure 20-3). Oral lesions are unusual in psoriasis; however, some authorities\(^\text{15-17}\) believe that the condition known as geographic tongue (Figure 20-4) may actually be a manifestation of psoriasis, because the histological features are identical. Occasionally, only intertriginous areas are involved (inverse psoriasis). When one suspects a diagnosis of psoriasis, it is often helpful to inspect the intergluteal cleft for involvement (Figure 20-5). A typical feature of psoriasis is Köebner’s phenomenon, the appearance of lesions in scars or other sites of trauma.\(^\text{18}\)

The lesions may appear in sites of old, major trauma such as surgical scars or in areas of recent or minimal injury such as an abrasion. The cause of this phenomenon is not known, although recently, speculation has focused on microbial factors.\(^\text{19}\)

The scale in psoriasis is typically silvery white and stacked in layers. In this way, the scale is said to be micaceous (slatelike). When the scale is removed with curettage or scraping, pinpoint bleeding may be noted. This is known as the Auspitz sign. It is more common, however, to observe a loose scale that easily flakes off in patients with long-standing lesions. Although traditionally con-
sidered a hallmark of psoriasis, the Auspitz sign’s sensitivity and specificity have been questioned by Bernhard, who elicited the sign in several nonpsoriatic, scaling disorders.20

When examining a patient with presumed psoriasis, one should always examine the fingernails and toenails, which frequently reveal diagnostic clues (Figure 20-6). Onycholysis—separation of the distal free edge of the nail plate from the nail bed—is a frequent finding. Less common, but more specific, is nail pitting—actual punctate depressions on the surface of the plate. The most specific nail finding, often said to be pathognomonic, is the so-called “oil spot,” which describes a yellowish brown, irregular macule beneath the nail plate and represents involvement of the nail bed with psoriasis.

Fig. 20-5. Annular erythematous plaque with fine surface scale in the intergluteal cleft. This is a common site for psoriasis.

Fig. 20-6. Psoriatic nails. (a) Pitting, thickening, and discoloration. (b) Pitting, distal onycholysis, and “oil spot.” (c) Distal onycholysis and “oil spots” in patient with large psoriatic plaques. (d) Onycholysis, discoloration, and crumbling of nail plate. Photograph c: Courtesy of Dr. Charles Trapp, MacDill Air Force Base, Fla.
sis. When the nail matrix is involved, severe onychodystrophy can result, with diffuse crumbling and yellowing of the plate. In pustular psoriasis and acrodermatitis continua of Hallopeau, one commonly observes subungual pustules.

Types of Psoriasis

Psoriasis Vulgaris

Psoriasis Vulgaris is the most frequent presentation of psoriasis, consisting of chronic, stable, well-defined plaques that may persist for years. While the plaques can occur on any cutaneous surface, they are most likely to appear in the areas of predilection described above.

Guttate Psoriasis

Guttate psoriasis most commonly occurs in young adults and is usually eruptive in onset. It consists of multiple, small, guttate (raindroplike) lesions over the trunk and proximal extremities (Figure 20-7). Frequently, careful history-taking will reveal an antecedent streptococcal pharyngitis or viral respiratory infection.21

Localized Pustular Psoriasis

Localized pustular psoriasis consists of two types: pustular psoriasis of the palms and soles, known as pustulosis palmoplantaris (Figure 20-8), and acrodermatitis continua of Hallopeau. Both forms present with pustules, the former involving the palms and soles, and the latter involving the distal fingers and toes with severe nail dystrophy (Figure 20-9). Both conditions are noted for chronicity and refractoriness to treatment. When severe, either can be disabling. Pustulosis palmoplantaris has been associated with arthritis of the anterior chest wall,22 cigarette smoking, and thyroid disease.23

Generalized Pustular Psoriasis

Generalized pustular (von Zambusch) psoriasis is one of the rarer presentations of psoriasis,24,25 marked by acute attacks of generalized pustules on erythematous skin and associated with fever, leukocytosis, and systemic toxicity. This disease can be fatal, so hospitalization is mandatory. Several provocative factors are claimed to precipitate attacks, but the most important to consider is the association with steroid withdrawal (ie, systemic and potent topical steroids).25,26

Psoriatic Arthritis

Psoriatic arthritis is an inflammatory, sero-negative arthritis clearly associated with psoriasis (Figure 20-10). It has been neatly classified into five general categories (Table 20-4).28

Diagnosis

When chronic, scaling plaques are present on extensor surfaces or on the penis or scalp, the diagnosis is usually obvious. Occasionally, however, psoriasis can mimic other papulosquamous disorders: seborrheic dermatitis, lichen planus, pityriasis rosea, pityriasis rubra pilaris, drug eruptions, or even syphilis. Isolated lesions may be confused with lichen simplex chronicus, nummular dermatitis, cutaneous lupus erythematosus, or mycosis fungoides. When one suspects the diagnosis of psoriasis, it is essential to perform a thorough cutaneous exam looking for clues such as nail involvement, evidence of Köebner’s phenomenon, or pinking of the intergluteal cleft. Obtaining a family history can also provide useful information. If doubt still exists, biopsy should be obtained because the histopathological features are fairly distinctive.

Treatment

Numerous modalities are available for the treatment of psoriasis. This fact speaks for the complexity of the disease, the mystery of its pathogenesis, and the limitations of all of the available treatments.
Fig. 20-8. (a) Psoriasis of the palms. (b) Psoriasis of the soles. (c) Pustulosis plantaris. (d) Close-up view of (c) to reveal characteristic yellow-brown pustules.
Common Skin Diseases

Fig. 20-9. Scaling and crusted lesions with border of peripheral pustules on the distal digits, characteristic of acrodermatitis continua of Hallopeau.

Fig. 20-10. Psoriatic arthritis. (a) Proximal interphalangeal joint of finger. (b) Toe involvement—"sausage digit." (c) Severe involvement of fingers and hand.
TABLE 20-4
ARThRITIS IN PSORIASIS

<table>
<thead>
<tr>
<th>Type of Arthritis</th>
<th>Characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asymmetric involvement of a few joints of the fingers</td>
<td>Affects 70% of patients with psoriatic arthritis; “sausage digit”</td>
</tr>
<tr>
<td>Symmetric polyarthritis</td>
<td>Affects 15% of patients with psoriatic arthritis; resembles rheumatoid arthritis</td>
</tr>
<tr>
<td>Distal interphalangeal joint involvement</td>
<td>Affects 5% of patients with psoriatic arthritis; “classic” psoriatic arthritis</td>
</tr>
<tr>
<td>Arthritis mutilans</td>
<td>Affects 5% of patients with psoriatic arthritis; deforming arthritis with bony destruction, telescoping of digits, and ankylosis</td>
</tr>
<tr>
<td>Ankylosing spondylitis</td>
<td>Affects 5% of patients with psoriatic arthritis; may also have peripheral joints involved</td>
</tr>
</tbody>
</table>


There clearly is no treatment of choice in the management of psoriasis. Many new modalities have been described recently, such as vitamin D, fish oil, and cyclosporine. At this point, these therapies should be considered investigational. Medical officers are advised to consider first the more traditional, time-honored modalities available.

No treatment is without side effects. The risks for adverse side effects can often be reduced by combination therapy, such as retinoids combined with psoralen and UV-A (PUVA). The advantage of combination therapy is that the cumulative dose of each agent is diminished. Thus, while the beneficial effects may be additive, the adverse effects are usually decreased. Furthermore, when approaching the treatment of psoriasis, one must realize that the aim of treatment should not be to obtain complete clearing. That is often an unrealistic approach in the management of psoriasis. Complete clearing is often unattainable without experiencing significant toxicity. Realistic expectations by the physician as well as the patient are perhaps the most important aspects of therapy.

**Topical Corticosteroids**

Topical corticosteroids are frequently employed in the management of psoriasis and can be quite beneficial. Due to a potent antiinflammatory effect and inhibition of deoxyribonucleic acid (DNA) synthesis (with a subsequent antiproliferative effect), topical steroids are a logical and effective modality in this disease. Intralvesional steroids may be considered for isolated or resistant plaques. The limiting factor for topical corticosteroids is epidermal and dermal atrophy, which can result in marked thinning of the skin and striae formation (Figure 20-11). More importantly, sustained use of potent topical steroids can lead to suppression of the hypothalamic-pituitary-adrenal axis. Pulse dosing may permit extended maintenance with potent topical steroids while limiting adverse effects. Another drawback to consider regarding steroid therapy is the development of tachyphylaxis. For these reasons, one is encouraged to consider alternative topical agents for the treatment of psoriasis.

![Fig. 20-11. Linear atrophic striae on the back of this patient, who is taking oral steroids.](image-url)
**Anthralin**

Anthralin is a synthetic anthrone derivative that reduces mitotic activity and suppresses free radicals. It is widely used in Europe as an integral component of the Ingram regimen, which consists of a daily coal tar bath followed by suberythemogenic UV-B exposure, then the application of dithranol paste of gradually increasing concentration. Although this method is quite effective and safe, it is cumbersome. For this reason, short-contact anthralin treatment plans have been devised, frequently with the addition of coal tar to diminish the irritancy of anthralin, which can limit its use. Due to its lack of systemic toxicity, carefully supervised anthralin therapy combined with thorough patient explanation and education is an ideal modality for the treatment of chronic plaque-type psoriasis.

**Coal Tar**

Coal tar has been widely employed in the treatment of psoriasis for more than a century. It is a complex mixture of over 10,000 substances, which may possess antimitotic activity. Coal tar gained popularity after introduction of a simple method described by Goeckerman in 1925, which consists of continued application of 2% crude coal tar, which is removed with mineral oil once daily for UV-B exposure. The tar is reapplied after bathing. Coal tar is an extremely safe agent. The most frequent side effect is a tar-induced folliculitis. There have been reports of skin cancers associated with coal tar application; however, this is an uncommon event, and such case reports usually involve patients who have been exposed to multiple other skin cancer–inducing agents.

**Systemic Methotrexate**

Methotrexate, given in weekly oral doses of 5 to 25 mg, is an extremely valuable agent for treating severe psoriasis and psoriatic arthritis that are not adequately controlled by more conventional therapies. It is a folate antagonist that exerts its antiproliferative effect on psoriatic epidermal cells by directly inhibiting DNA synthesis. Methotrexate is absolutely contraindicated in pregnant or lactating females. Relative contraindications are many, including liver or renal abnormalities, excessive alcohol consumption, and anemia, leukopenia, and thrombocytopenia. Guidelines for appropriate use of methotrexate in psoriasis are published periodically by the Psoriasis Task Force of the American Academy of Dermatology. Medical officers should be thoroughly knowledgeable with these guidelines before considering methotrexate for the treatment of psoriasis. The greatest limitation of methotrexate is its hepatotoxicity. For this reason, newer folate antagonists with perhaps fewer effects on the liver are being investigated.

**Systemic Retinoids**

The retinoids are synthetic derivatives of vitamin A and vitamin A acid. The effects of vitamin A and vitamin A analogs on the growth and terminal differentiation of epidermal cells and the vitamin’s beneficial effect on disorders of keratinization have long been known. Retinoids have been shown to affect growth of epidermal cells by altering keratin synthesis and formation of the cell envelope. Etretinate (0.5–1.0 mg/kg/d) is the retinoid approved for use in the treatment of psoriasis. It should be considered for initial treatment in the management of pustular psoriasis or erythrodermic psoriasis. It is also beneficial for the patient with extensive plaques and severe involvement recalcitrant to more conventional modalities and has been successful in clearing acrodermatitis continua of Hallopeau.

Etretinate has significant risk for adverse effects including teratogenicity, elevation of serum lipids, liver toxicity, and numerous mucocutaneous side effects. In addition, cases of skeletal hyperostoses have been documented with long-term use. Furthermore, it is known to be stored in adipose tissue for long periods, thus leading to a prolonged half-life (> 100 d). In fact, blood levels are detectable for more than 2 years after discontinuation of therapy. For these reasons, in addition to its potent teratogenicity, etretinate is best avoided in women of child-bearing potential. Alternatives include isotretinoin and, once available, acitretin—the main metabolite of etretinate—which has been shown to be effective in treating psoriasis and has a much shorter half-life (50 h for acitretin vs 120 d for etretinate). Because the side effects of acitretin are similar to those of etretinate, including potent teratogenicity, effective contraception is essential for women and long-term use is discouraged. Because risks are greatly increased with long-term use of retinoids, they are probably best used in combination with other modalities such as phototherapy.
**Ultraviolet-B Phototherapy**

Natural sunlight has long been known to improve psoriasis. An excellent (and more reliable) substitute is an artificial light source that produces light primarily in the ultraviolet-B (UV-B) range (290–320 nm). This allows for continuous monitoring and metered delivery of UV radiation, thus diminishing the potential for accidental overexposure and burning. Although the mechanism of the therapeutic effect is unknown, UV-B alone or with coal tar is an excellent method for clearing psoriasis and maintaining remission. Its use should be considered early in the course of management.

**Psoralen and Ultraviolet-A Phototherapy**

Introduced in 1974 by Parrish and Fitzpatrick, PUVA phototherapy has gained worldwide acceptance in the treatment of psoriasis. Patients ingest a specific dose of methoxsalen (consult chart in package insert for precise dosage based on weight), followed in approximately 90 minutes by a carefully metered UV-A (320–400 nm) exposure, which is increased with progressive treatments. Initially, Oxsoralen was the drug employed. This has now been replaced by Oxsoralen-Ultra (methoxsalen, manufactured by ICN Pharmaceuticals, Inc., Costa Mesa, Calif.), a liquid form in a gel capsule that provides more uniform serum drug levels. Patients are treated two to three times per week, and generally respond within 20 to 25 treatments. PUVA can then be continued and gradually tapered to maintain prolonged remission.

PUVA exerts its effect through the production of bifunctional DNA adducts and subsequent inhibition of DNA synthesis and epidermal cell replication. It is an excellent modality for managing severe psoriasis on an outpatient basis. However, it is not without risk. Patients must wear protective eyewear (wraparound UV-A filtering glasses) for 24 hours after ingestion of the drug to prevent cataracts. It has been well documented that PUVA is clearly associated with an increased risk for nonmelanoma skin cancer, particularly squamous cell carcinoma. The study by Fitzpatrick and Parrish reveals that the risk of developing squamous cell carcinoma increases with cumulative PUVA exposure. The higher exposure group (> 260 treatments) has a risk that is 12-fold that of the lower exposure group (< 160 treatments) and 100-fold that of the general population. This fact emphasizes the point made previously: the goal of therapy when managing psoriasis should not be 100% clearing. Ninety percent clearing is, generally, much easier to attain and maintain at far less risk, and this is usually quite acceptable to the patient. Clearly, PUVA should not be used unless the patient’s psoriasis is severe. While PUVA-induced squamous cell carcinoma has little potential for metastatic spread, all patients receiving PUVA must be carefully monitored to ensure early detection and prevention of disfigurement. All men should shield the genital region during therapy, as this area has proven to be at particular risk for developing PUVA-induced tumors.

PUVA-associated carcinogenesis may possibly be avoided by employing PUVA bath treatments using a trioxsalen solution. This technique has been used extensively in Europe and has been shown to be of equal efficacy to and probably safer than standard PUVA. This author has found a similar technique using a solution of Oxsoralen, as described by Coleman et al, to be extremely effective in the management of pustulosis palmpoplantaris. All forms of PUVA may be combined with other treatments such as retinoids to obtain superior results while diminishing adverse effects.

**Military Considerations**

The main point to emphasize to the military physician regarding psoriasis is its tendency towards chronicity and its sometimes unpredictable course. Consider the individual with mild plaque-type psoriasis who is easily controlled with anthralin and intermittent UV-B therapy. That same individual deployed to a northern climate or hostile environment may experience a significant flare-up requiring medical evacuation and preventing mission completion. This author has observed this scenario repeatedly. In World War II, dermatologists found that approximately 20% of psoriatic patients became liabilities and questioned whether troops with significant psoriasis should ever be sent overseas. In U.S. Army medical clinics in Vietnam, psoriasis accounted for up to 6% of outpatient visits. Individuals with psoriasis of even mild severity should be considered for presentation to a medical evaluation board for determination of worldwide qualification.

Another point to consider about psoriasis in the military is a controversial one: the role of antimalarial chemoprophylaxis in inducing exacerbations of psoriasis. This author witnessed one case of a young sailor who experienced a pustular flare-up of his psoriasis while on chloroquin prophylaxis. This required medical evacuation from a
ship and a prolonged hospital course, including an initial stay in the intensive care unit. While this is anecdotal, it illustrates the argument for reconsidering worldwide eligibility for all military members with psoriasis. Kuflik reported on 48 U.S. Army patients with psoriasis on prophylactic chloroquin in Vietnam: 42% of the cases worsened, but only 6% had significant flare-ups.64

LICHEN PLANUS

Lichen planus (LP) is an inflammatory skin disease that is often pruritic, with distinctive mucocutaneous findings. In 1969, Erasmus Wilson published a series of 50 cases describing the classic features of LP.65 The prevalence of LP in the general population is estimated to be 0.13% to 0.34% for cutaneous involvement and 0.19% to 1.5% for oral involvement.66 In a study of over 670,000 patients in the United States, LP was seen in 0.44%.67 In a general dentistry clinic, oral LP was seen in 0.6%.68

LP predominantly affects people in their middle ages, with approximately two thirds of those affected between the ages of 30 and 60.67 It is worldwide in its distribution with no racial predisposition. Both sexes are affected approximately equally. Familial cases have been reported.69,70

Clinical Features

The characteristic lesion of LP is a flat-topped, shiny, polygonal papule that is most often described as violaceous in color. Individual papules range in size from pinpoint to greater than 1 cm and may remain discrete or become confluent. Frequently, one can observe fine, reticulate white streaks on the surface. These streaks, known as Wickham’s striae, are most easily seen with the use of mineral oil and

Fig. 20-12. Lichen planus. (a) Ankle. Note violaceous, flat papules with reticulated white scale. (b) Flexor surface of the wrist, a common location. (c) Dorsum of hand. (d) Close-up of (a) to demonstrate Wickham’s striae.
the aid of a hand-held lens. Köebner’s phenomenon is sometimes observed.18,71

LP may remain a localized process or have a generalized distribution. When LP becomes generalized it usually does so within the first month of onset and is typically symmetric in its distribution. The majority of cases begin on the extremities, especially the ankles72 and the flexor aspects of the wrists (Figure 20-12).73 The ankles and shins are the most common sites for hypertrophic lesions.74 Other cutaneous sites commonly involved include the lumbar region and the penis; the latter site may have annular or ulcerative lesions (Figure 20-13). Erosive and ulcerative forms of LP can also be found on other mucous membranes as well as on the palms, soles, and arms (Figure 20-14).75,76

The natural course of LP is usually spontaneous resolution—93% within 2 years in Samman’s study.77 Among this same series of patients, the rate of relapse was 20%. Typically, marked post-inflammatory hyperpigmentation, hypopigmentation, or both will occur as lesions regress.78

Nail involvement is seen in 10% of cases.77 This is most commonly manifested as surface roughness or flaking with longitudinal ridging. Other changes include brownish discoloration, pitting, and thinning of the nail plate with splitting of the free edge. Chronic nail involvement can result in pterygium formation as a result of the growth of the cuticle and subsequent adherence to the nail bed, resulting in permanent loss of the nail plate. Occasionally, the nails may be the only site of involvement.79,80 These cases normally require histological confirmation for diagnosis. Twenty-nail dystrophy, a disorder of acquired dystrophic nail changes in children, may be a manifestation of LP.81,82 Several other variants of cutaneous LP exist (Exhibit 20-1).

Involvement of oral mucosa is very common in LP. In Altman and Perry’s series of 197 patients, 40% had cutaneous and mucosal involvement, and 25% had oral alone.83 Oral lesions can be found in up to 71% of LP patients seen in dermatology clinics.77,83 Oral LP typically involves buccal mucosa, although other sites can be involved. These include

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**EXHIBIT 20-1**

**VARIANTS OF CUTANEOUS LICHEN PLANUS**

- Hypertrophic lichen planus
- Linear lichen planus
- Annular lichen planus
- Atrophic lichen planus
- Lichen planus subtropicus (actinicus)
- Bullous lichen planus
- Lichen planopilaris
- Lichen planus pemphigoides
- Lichen planus–lupus erythematosus overlap
- Ulcerative lichen planus
- Hepatitis-associated lichen planus
**Common Skin Diseases**

**Fig. 20-15.** Oral lichen planus. (a) Characteristic reticulate, white plaque on the buccal mucosa—the most commonly involved site. (b) Involvement of the lower lip.

The tongue, gingiva, and vermilion border of the lip (Figure 20-15). Involvement of the palate and sublingual region is rare. Almost always, oral LP occurs bilaterally. The appearance of LP in the oral mucosa can be quite varied. The most common morphology is that of a plaque consisting of slightly elevated, very fine white lines in a reticulate pattern, similar to Wickham’s striae. This has been referred to as “Honniton lace.” Other forms include a papular variant or a plaquelike form (resembling leukoplakia), an atrophic form (resembling erythroplakia), bullae (rare), and an erosive form. Erosions are often extensive and can result in desquamative gingivitis.

The natural course of oral LP is one of chronicity, with a mean duration of almost 5 years. Oral LP has been shown to persist for more than 25 years. Persistent, erosive, oral LP may have a potential for the development of squamous cell carcinoma, although this is controversial. Some authorities consider oral LP to be a premalignant condition.

**Pathogenesis**

The precise etiology of LP remains unknown despite intensive investigation. Multiple hypotheses support numerous etiologies, including infectious agents, tobacco, betel nuts, psychogenic stress, chemicals, and numerous drugs. Some cases of LP have occurred in patients with diabetes, hepatitis, hepatic biliary cirrhosis, hypertension, and urolithiasis. While the exact triggering mechanism for LP remains unknown, recent work strongly implicates T-cell-mediated cytotoxicity as the final common pathway leading to the development of the lesion of LP. Norris recently demonstrated marked staining for ICAM-1 (intercellular adhesion molecule-1) of basal cells in skin from lesions of lichen planus. One could propose that this would lead to binding between the basal keratinocytes and T lymphocytes within the dense dermal infiltrate, resulting in the cytotoxic destruction of the keratinocytes.

**Diagnosis**

Lichen planus has a rather distinctive clinical presentation. Other entities to consider in the differential diagnosis are lichenoid drug eruptions, psoriasis, syphilis, and LP-like keratoses. When the skin lesions suggest a diagnosis of LP, always examine the mouth, which will frequently reveal the characteristic white, reticulate plaques typical of LP. Oral lesions, in the absence of cutaneous findings, may be confused with candida, traumatic erosions, aphthae, leukoplakia, or bullous diseases such as pemphigus vulgaris or erythema multiforme. Biopsy will usually confirm the diagnosis of LP, which has specific histological features. History is the most important feature to distinguish a lichenoid drug eruption from LP.

**Treatment**

The mainstay of therapy for LP is corticosteroids. In localized forms, potent topical steroids or...
steroids applied under occlusion may be beneficial. Large plaques generally respond to intralesional triamcinolone (3–5 mg/mL). Generalized or eruptive LP usually requires systemic steroid therapy. Alternatives to consider are PUVA95,96; retinoids97,98; and griseofulvin (125–250 mg twice daily), which has been reported to have a cure rate as high as 90%,99,100 although other reports show it to be completely ineffective.101 Oral lesions may respond to topical steroids in Orabase (gelatin, pectin; manufactured by Colgate-Hoyt, Canton, Mass.) or aerosolized betamethasone valerate. Recently, two groups102,103 have reported significant improvement of oral LP using a topical cyclosporine rinse with no systemic adverse effects and little systemic absorption over a 2-month period. However, the cost of cyclosporine may prove prohibitive.

In addition, antihistamines and antipruritic lotions may be useful and should be employed for relief of pruritus.

Military Considerations

Although LP is generally considered to be a benign, self-limited disease, the pruritus that usually accompanies it may interfere with performance of duty. It is important, then, to treat symptomatic patients aggressively to keep them functional.

Of historical significance is the widely reported relationship of atypical LP with the administration of quinacrine for antimalarial prophylaxis in World War II.104–107 The study by Bazemore et al104 of 400 patients reveals that long-term therapy with antimalarials is necessary to induce atypical LP, as only 20% of patients developed the disease within 3 months, while 80% developed the disease within 7 months. Many cases were complicated by a secondary pyoderma—usually caused by Streptococcus pyogenes and Staphylococcus aureus. This most likely was a result of inadequate hygiene in a tropical environment. The eczematous process frequently involved the hands, thus disqualifying many individuals from full military duty. On return to duty, relapse was almost certain, so prompt evacuation became the treatment of choice.62

A variant of LP rarely seen in the United States that could potentially be of significance for deployed military personnel is LP subtropicus (actinicus). This is common in the Middle East and consists of annular lesions on sun-exposed surfaces, particularly the face.108–110 The course is chronic and the condition worsens with sun exposure. Treatment consists of using sun blockers and limiting sun exposure.

ATOPIC DERMATITIS

Atopic dermatitis is an environmentally induced disorder occurring in genetically predisposed individuals. It is characterized by acute episodes of eczematous cutaneous eruptions with characteristic distribution, usually accompanied by xerosis. The disorder may be considered part of the atopic diathesis that also includes allergic rhinitis, conjunctivitis, and asthma. It is prevalent worldwide and is more common among children, particularly of industrialized countries. Its incidence in England is 1.1% to 3.1%.111,112 In the United States it is 0.7 to 2.4%.113

Clinical Features

The clinical features of atopic dermatitis can be classified into major features that are seen in most atopic patients and minor, nonspecific features that are frequently seen in atopic patients (Exhibit 20-2).114 There is no primary lesion of atopic dermatitis. The lesions are eczematous and marked by erythema, weeping, scaling, crusting, and lichenification (thickening of skin in response to continual rubbing). Often, the lesions may be excoriated or secondarily infected. Atopic dermatitis is recognized by observing these findings in a typical pattern of distribution. In infants and children under 2 years of age, the lesions typically occur on the face (Figure 20-16) and extensor surfaces. Paradoxically, in older children and adults, the characteristic pattern is that of predominantly flexural involvement with sparing of the face (Figure 20-17). History is extremely helpful in establishing the diagnosis of atopic dermatitis. The age of onset is usually in childhood—60% within the first year of life.115 Seventy percent of patients will have a history of asthma, hay fever, or both.114 A family history of atopy is almost always present.
### EXHIBIT 20-2

**CLINICAL FEATURES OF ATOPIC DERMATITIS**

<table>
<thead>
<tr>
<th>Major Features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pruritus</td>
</tr>
<tr>
<td>Flexural lichenification in adults</td>
</tr>
<tr>
<td>Facial and extensor involvement in infants and children</td>
</tr>
<tr>
<td>Chronic or chronically relapsing dermatitis</td>
</tr>
<tr>
<td>Personal or family history of atopy (asthma, allergic rhinitis, atopic dermatitis)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Minor Features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Xerosis</td>
</tr>
<tr>
<td>Ichthyosis, palmar hyperlinearity, keratosis pilaris</td>
</tr>
<tr>
<td>Immediate (type 1) skin test reactivity</td>
</tr>
<tr>
<td>Elevated serum immunoglobulin E</td>
</tr>
<tr>
<td>Early age of onset</td>
</tr>
<tr>
<td>Tendency toward cutaneous infections (especially with <em>Staphylococcus aureus</em> and herpes simplex virus), impaired cell-mediated immunity</td>
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<tr>
<td>Tendency toward nonspecific hand or foot dermatitis</td>
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<tr>
<td>Nipple eczema</td>
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<tr>
<td>Cheilitis</td>
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<tr>
<td>Recurrent conjunctivitis</td>
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<tr>
<td>Dennie-Morgan infraorbital fold</td>
</tr>
<tr>
<td>Keratoconus</td>
</tr>
<tr>
<td>Anterior subcapsular cataracts</td>
</tr>
<tr>
<td>Orbital darkening</td>
</tr>
<tr>
<td>Facial pallor, facial dermatitis</td>
</tr>
<tr>
<td>Pityriasis alba</td>
</tr>
<tr>
<td>Anterior neck folds</td>
</tr>
<tr>
<td>Pruritus when sweating</td>
</tr>
<tr>
<td>Intolerance to wool</td>
</tr>
<tr>
<td>Perifollicular accentuation</td>
</tr>
<tr>
<td>Food intolerance</td>
</tr>
<tr>
<td>Course influenced by environment or emotional factors</td>
</tr>
<tr>
<td>White dermographism, delayed blanch</td>
</tr>
</tbody>
</table>


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**Fig. 20-16.** Atopic dermatitis in an infant most commonly involves the face, with perioral sparing.

**Fig. 20-17.** Atopic dermatitis in a young adult, with typical involvement of antecubital fossae and marked symmetry.

**Natural Course**

The natural course of atopic dermatitis is one of repeated episodes of acute flare-ups, followed by a subacute phase of slow resolution. Acute episodes tend to occur in response to psychic or emotional stress. In a study in which more than 500 patients were interviewed, stress was considered the primary precipitating factor. Between flare-ups, patients will often be bothered by generalized xerosis, which often is quite pruritic and creates an intolerance of the skin for exposure to solvents, soaps and other cleansers, and water (in excessive amounts).
Hand dermatitis is frequently seen in patients with atopic dermatitis, particularly adults. Approximately 70% will experience it. Careful questioning of adult patients with hand dermatitis will often reveal a prior history of atopic dermatitis.

Complications

Staphylococcal pyoderma is a frequent complication of atopic dermatitis. The skin of nearly all patients with atopic dermatitis is heavily colonized with \textit{S. aureus}. This is partially explained by the observation that \textit{S. aureus} may have specific binding sites for tissue fibronectin and laminin. In general, these infections remain superficial, but cases of osteomyelitis have been reported in children with severe atopic dermatitis.

As a result of impaired cellular immune responses, patients with atopic dermatitis sometimes are unable to adequately handle viral or fungal infections of the skin. This increased susceptibility has resulted in widespread cutaneous viral infections such as eczema vaccinatum (vaccinia virus from smallpox vaccine) and Kaposi’s varicelliform eruption, which is usually caused by herpes simplex virus (HSV) (Figure 20-18), but has also been seen with coxsackievirus A16. Both produce a similar clinical picture of an acute, widespread, vesiculopustular eruption marked by fever, adenopathy, and prostration. The patients may appear quite ill and some have died. Treatment is with intravenous acyclovir (750 mg/m$^2$/d) and antibiotics for secondary bacterial infection. Other viruses that can cause problems for atopic patients include molluscum contagiosum and human papillomavirus (HPV).

Diagnosis

Atopic dermatitis is most often confused with contact dermatitis (irritant or allergic), but other dermatoses in the differential diagnosis include seborrheic dermatitis, psoriasis, scabies, nummular dermatitis, and lichen simplex chronicus. Atopic dermatitis is a clinical diagnosis based on the findings of at least three major and three minor criteria (see Exhibit 20-2).

Treatment

When approaching the management of atopic dermatitis, it is useful to keep in mind that this is a disease that is precipitated in genetically predisposed individuals by an environmental stress. The first goal of treatment, therefore, is the elimination of known precipitating factors (Exhibit 20-3).

Topical steroids are a mainstay of therapy in relieving the inflammation of acute episodes and the associated pruritus. Their use should be carefully supervised by the treating physician. Patients must be aware of the risks for dermal atrophy, adrenal suppression, and tachyphylaxis. Because flexural areas are commonly involved, one must be especially alert for possible steroid-induced atrophy. Topical steroids should be applied sparingly and only on inflamed areas. Highly potent topical steroids such as clobetasol propionate applied twice daily may be useful in short courses (less than 2 wk) for treating areas of lichenification, but are best avoided in children. Systemic steroids, while effec-

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**EXHIBIT 20-3**

**COMMON PRECIPITATING FACTORS IN ATOPIC DERMATITIS**

- Excessive exposure to soap, hot water, or chemicals
- Psychic stress
- Overheating (by physical exertion or hot climate)
- Cutaneous infection or infestation (eg, scabies)
- Extremes of temperature (hot or cold)
- Prickly clothes
- Allergic exposure (when relevant)

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Fig. 20-18. Kaposi’s varicelliform eruption from herpes simplex virus in a young adult female with a history of eczema in childhood.
ative, are not recommended as patients are likely to become dependent on them.

Because xerosis is so often a precipitating factor, hydration of the skin is an essential component of therapy. Emollients such as Eucerin (water, mineral oil; manufactured by Beiersdorf, Norwalk, Conn.), Lac-Hydrin (ammonium lactate, manufactured by Westwood Pharmaceuticals, Buffalo, N.Y.), Moisturel (petrolatum, glycerin; manufactured by Westwood Pharmaceuticals, Buffalo, N.Y.), and 10% urea cream are effective agents that are best applied after bathing. They can be applied to the entire cutaneous surface; topical steroid is applied only to inflamed areas.

Antihistamines such as hydroxyzine (25 mg three times daily), cyproheptadine hydrochloride (4 mg three times daily), and doxepin (75–150 mg/d) can be useful for relieving pruritus. They may be as useful for their associated antiserotonin, anticholinergic, anxiolytic, and sedative effects as for their antihistaminic effect. Topical antipruritic lotions such as Sarna lotion (camphor, menthol, phenol; manufactured by Stiefel, Coral Gables, Fla.) may also be helpful.

Antibiotics are often useful in the management of acute episodes as secondary impetiginization is very common. Erythromycin and dicloxacillin (for both drugs, 250–500 mg twice daily for 10 d) are useful agents and topical mupirocin (applied topically twice daily for 10 d) has been shown to be effective.19

Coal tar may be a useful adjunct to therapy. Liquor Carbonis Detergens (coal tar, nonproprietary) can be compounded with a topical steroid such as triamcinolone acetonide. This particular combination can be quite effective.

Phototherapy can be very beneficial in the management of atopic dermatitis. Successful results have been reported with PUVA,127 UV-B,128 and a combination of UV-A and UV-B.129 As these modalities command significant patient cooperation and sophistication, and have long-term risks, they are not recommended for preadolescents. Because PUVA has the greatest long-term risks, it should be reserved for severe, refractory cases.

Military Considerations

One need only glance at the common precipitating factors (see Exhibit 20-3) to realize that life in the military can be hostile for the individual with atopic dermatitis. It is difficult, if not impossible, to eliminate all of these factors from the military environment. It is not surprising, then, that Sulzberger estimated that atopic dermatitis accounted for 3% to 5% of all disease-related military discharges during World War II.133 It was the opinion of most dermatologists who served in World War II that it was not prudent to send overseas any man with a history of atopic dermatitis, as these men frequently required prolonged and repeated hospitalization.62

Frequently, an individual will appear to have “outgrown” his or her atopic dermatitis and, indeed, will not have experienced an outbreak for several years. It is important to remember, however, two features of the atopic diathesis: a propensity for hand dermatitis, and an impaired cellular immune response. Both factors are negative considerations for a military career.

Hand dermatitis is most likely to result from exposure to chemical irritants. Often, the exposure is occupationally related. Studies have determined that of patients with an occupational dermatitis, 85% have a personal or family history of atopic disease.134,135 An individual with a history of atopic dermatitis and a propensity for developing hand dermatitis would be disqualified from many mili-
tary occupations in order to avoid duty-related exposure to irritant chemicals.

Impaired cellular immunity renders the atopic patient susceptible to disseminated vaccinia infection (eczema vaccinatum). For this reason, smallpox vaccination becomes dangerous, even if the atopic dermatitis is in remission. Although smallpox vaccination is no longer recommended, the military could require it under special circumstances.

**URTICARIA**

Urticaria is a common pruritic eruption, marked by characteristic evanescent lesions known as wheals. It affects approximately 15% to 20% of the population at some point during their lifetime.\(^{136}\) Wheals are, by definition, superficial in nature. When the edematous process spreads to involve the deep dermis and subcutaneous tissue, then it is termed angioedema (Figure 20-19). The edema can involve other organ systems, particularly the cardio pulmonary system and the gastrointestinal tract. Patients can present with urticaria alone (40%), angioedema alone (10%), or urticaria and angioedema combined (50%).\(^{137}\)

**Clinical Features**

The clinical hallmark of urticaria is the wheal or hive. It is an elevated, sharply demarcated, pale red or white, dermal flat-topped plaque with no associated epidermal change. Wheals may be bordered by a thin, pale, or red halo (Figure 20-20). Their size varies from a few millimeters to several centimeters and they may be round or oval or coalesce into large, bizarre polycyclic or serpiginous forms. The most characteristic feature of the wheal is its evanescent nature. Although the process can last weeks or even months, individual lesions usually resolve in a matter of hours and rarely persist beyond 24 hours. Wheals are typically quite pruritic and can occur anywhere on the integument.

The occurrence of systemic symptoms is not unusual in urticaria or angioedema. For this reason, a thorough history—including an extensive review of systems—and a physical examination are required. Associated symptoms include arthralgias, headache, hypotension, hoarseness, dyspnea, abdominal pain, nausea, and diarrhea. Because angioedema is a deeper process that more commonly involves mucosal surfaces, it is more likely...
to cause these systemic symptoms, which can be life-threatening.

Urticaria has an unpredictable course. The majority of cases are acute and self-limited with spontaneous resolution after a few hours to several days. Many of these patients do not seek medical attention. Urticaria that persists beyond 8 to 12 weeks is arbitrarily classified as chronic urticaria. Approximately 50% of patients are lesion-free in less than 1 year, while 20% may experience recurrent episodes beyond 20 years.138

Types of Urticaria

The urticarias can be grouped in a variety of clinical entities. Champion has described some 50 separate classifications.139,140 Some of the more important types of urticaria are discussed below.

Physical Urticarias

The physical urticarias account for approximately 16% of all urticarias seen in specialty clinics.139 Over 20 types have been described. Understanding physical urticaria will permit the identification and possible elimination of the etiologic mechanism, thus preventing recurrent episodes.

Dermatographism is the most common of the physical urticarias (Figure 20-21). It is easily reproducible by the firm, brisk stroking of the skin with a blunt-tipped object. The resultant wheal occurs in minutes in the exact location that the pressure was applied. It can be elicited in approximately 1.5% to 4% of healthy individuals.141

Pressure urticaria is a much less common form, characterized by deep, sometimes painful, wheals that occur at sites exposed to deep pressure for prolonged periods (Figure 20-22)—usually the buttocks or feet. The timing of these wheals is unique in that they occur hours after pressure has been applied and often persist for more than 24 hours. In a study of 44 patients with pressure urticaria, Daves et al142 determined that the mean onset of whealing was 3.5 hours with a peak at 10 hours, and a mean lesion duration of 36 hours. In their experience, systemic corticosteroids were the only effective treatment. Fifty percent of the patients also exhibited delayed dermatographism. Delayed dermatographism has also been described143 as an independent entity, distinct from pressure urticaria. In general, however, dermatographism is easily distinguished from pressure urticaria by its immediate onset (minutes versus hours).

Cold urticaria represents from 1% to 7% of all forms of the physical urticarias.144,145 It occurs in an acquired and a familial form with autosomal dominant inheritance. The primary acquired form is the most common (96%),146 and the familial form is quite rare.147 Cold urticaria has been associated with cryoglobulins, cryofibrinogens, cold agglutinins, and paroxysmal hemoglobinuria.146 Such cases are referred to as secondary acquired cold urticaria. In response to cold stimuli, localized symptoms are produced (erythema, pruritus, wheals), as well as systemic manifestations including respiratory distress, hypotension, and even anaphylaxis. The di-

Fig. 20-22. Pressure urticaria. Wheals occurred when the patient lay on his back for prolonged periods. The circled wheals occurred at the exact location where pressure had been previously applied with a ballpoint pen.
agnosis is easily confirmed by applying an ice cube to the skin. A wheal will occur several minutes later during the rewarming phase. The ice cube test is negative in the familial type. The duration of cold urticaria is from 3 weeks to over 37 years. Patients are advised to avoid cold climates and rapid decreases in temperature, as may occur in ocean swimming.

Solar urticaria is a rare condition characterized by wheal formation within minutes of sun exposure, sometimes with associated systemic symptoms. Lesions can be induced by light of varying wavelengths from the UV-B range (290–320 nm) to the visible light spectrum (> 400 nm). Although the precise mechanisms of lesion induction are not entirely clear, Leiferman et al provided convincing evidence for the role of eosinophil degranulation in the pathogenesis. In addition, a circulating photoallergen has been described that is generated by absorption of light energy. Removal of this factor by plasmapheresis has been shown to result in prolonged remission. Tolerance may be induced by PUVA or UV-A alone.

Aquagenic urticaria is precipitated by contact with water, regardless of temperature. It is a rare condition that resembles cholinergic urticaria (described below). Shelly and Rawasley reported successful control with ChlorTrimeton (chlorpheneramine maleate, manufactured by Schering Corp., Kenilworth, N.J.), 4 mg four times daily. Aquagenic urticaria is a separate clinical entity from aquagenic pruritus.

Vibratory angioedema is also a rare condition in which localized erythema and edema occur in response to vibration. Autosomal dominance has been described, as well as an acquired form that can be occupationally related. A good therapeutic response to terfenadine (60 mg twice daily) has been reported in at least one case.

Localized heat urticaria, another rare physical urticaria, occurs in response to skin exposure to heat above 43°C. Successful induction of tolerance through graduated incremental exposure under medical supervision has been reported.

Cholinergic Urticaria

In cholinergic urticaria, which accounts for 4% of urticarias, wheals are provoked by heat, emotion, or gustatory stimuli. The lesions produced are characteristically tiny (1–3 mm), markedly pruritic, and persist for roughly 45 minutes to 1 hour. The cutaneous response is felt to be produced by the action of acetylcholine on the mast cell. A new entity of exercise-induced anaphylaxis that is distinguishable from cholinergic urticaria has been described. Cholinergic urticaria is usually easily treated with antihistamines.

Contact Urticaria

Contact urticaria is an uncommon condition that consists of a wheal-and-flare response to a variety of substances applied to the skin, and occurs within 20 to 30 minutes after application. Anaphylaxis can result; it has been described in 12 individuals during surgery, and was caused by their hypersensitivity to natural latex in the surgical glove.

Urticarial Vasculitis

Leukocytoclastic vasculitis presenting as urticaria or angioedema was first recognized at the Mayo Clinic. The cutaneous findings may be identical to common urticaria except that the wheals may be somewhat purpuric, tend to persist for 3 to 5 days, and often heal with residual hyperpigmentation. It is an important entity to recognize because of its frequent association with systemic diseases such as serum sickness, systemic lupus erythematosus, Sjögren’s syndrome, and infections such as viral hepatitis type B.

Hereditary Angioedema

Hereditary angioedema is a rare cause of angioedema with its onset in childhood and autosomal dominant inheritance. The episodes of swelling usually recur frequently, are often painful and persistent, and are aggravated by trauma. This form of angioedema occurs without urticaria, so the presence of wheals should exclude the diagnosis. The angioedema is due to a lack of functional C1 esterase inhibitor, an enzyme that controls complement activation. The inhibitor may be diminished in quantity, or may be present but dysfunctional. Daily administration of danazol, an anabolic steroid, may prevent attacks.

Urticaria due to Histamine-Releasing Agents

Many substances (Exhibit 20-4) are capable of causing direct mast cell degranulation with subsequent histamine release. This is not an immune-modulated mechanism. The resultant wheals occur shortly after exposure. Aspirin may exacerbate
Common Skin Diseases

That have an obvious cause will be self-evident, so those patients will often not seek medical attention. The process may be multifactorial, so careful history taking is mandatory to try to uncover possible precipitating factors (Exhibit 20-5).

Diagnosis

It is tempting to order multiple screening laboratory tests to rule out the rarer causes of urticaria. Although it may be reasonable to order a few tests, such as a complete blood count or an erythrocyte sedimentation rate, extensive laboratory investigation is often pointless and expensive, and should not be routine. What is important and should be routine is a thorough history and physical examination. The history should be directed at determining whether or not a definable type of urticaria exists that may be related to a physical stimulus, an inhalant, an ingested substance, a systemic disease, an infection, or emotional stress. The information obtained from both the history and the physical will guide any laboratory evaluation. If no specific cause is found, and the symptoms warrant, the patient should be treated empirically.

Treatment

The majority of patients with idiopathic urticaria will respond to antihistamine therapy. H$_1$-receptor antagonists are the major class of therapeutic agents employed in the management of urticaria. Atarax (hydroxyzine, manufactured by Roerig, New York, N.Y.), 25 mg three times daily, is an excellent first-line therapy and dose may be increased to tolerance, with sedation being its primary dose-limiting effect. It is often useful to add a second H$_1$ blocker from a different chemical class. Combination with an H$_2$-receptor blocker such as cimetidine (300 mg four times daily) may also be helpful. Similarly, good results have been obtained with doxepin (75–150 mg/d), an antidepressant with known anti-H$_1$ and anti-H$_2$ activity, in the management of chronic idiopathic urticaria.

Because of the problems with sedation from the traditional antihistamines available, much effort went into the development of newer agents with fewer central nervous system (CNS) side effects. Of these second-generation antihistamines, Hismanal (astemizole, manufactured by Janssen Pharmaceuticals, Piscataway, N.J.), 10 mg daily, is the only one currently approved for the treatment of urticaria, but Seldane (terfenadine, manufactured by Marion
### EXHIBIT 20-5
### CAUSES OF URTICARIA

<table>
<thead>
<tr>
<th>Foods</th>
<th>Drugs</th>
<th>Systemic Diseases</th>
<th>Other</th>
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<td>Shellfish</td>
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<td>Eggs</td>
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<td>Systemic lupus erythematosus</td>
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Merrell Dow, Kansas City, Mo.), 60 mg twice daily, can also be beneficial.

### Military Considerations

Because most cases of urticaria resolve spontaneously in a short period of time, they usually do not become a problem of any special significance in the military. Patients with a physical urticaria could have military occupational exposures that would exacerbate the disease. This obviously poses a potential threat to their health and safety, or to the safety of others, as well as to mission accomplishment. It is hard to imagine a career in the military, particularly in time of war, protected from physical stresses such as heat, cold, pressure, or vibration. Although comprehensive etiologic studies of urticaria and angioedema are not available from World War II, hospital admission statistics do reveal that there were almost 30,000 admissions for urticaria and more than 7,000 for angioedema between 1942 and 1945. In dermatology clinics in Vietnam, urticaria was seen in 1.75% to 2.74% of patients examined.

Another aspect of urticaria that the military physician must consider is the potential harm from the side effects of therapy. As discussed, antihistamines, the mainstay of therapy for urticaria, can have profound CNS side effects—particularly sedation. Again, this could jeopardize individual safety and mission accomplishment. Thus, the physician must address each patient’s military occupation and consider imposing temporary duty restrictions. The newer antihistamines, by virtue of their diminished permeability of the blood–brain barrier, will lessen the hazards.
Pseudofolliculitis barbae (PFB) is a condition that occurs in black men in response to shaving. It is common in the military services, where shaving is mandatory, with the prevalence estimated to be 45%\(^{170}\) to greater than 80%.\(^{171}\)

**Clinical Features**

PFB typically manifests as perifollicular, inflammatory papules and papulopustules of the beard area, particularly the anterior portion of the neck and the submandibular region. On close inspection, ingrown hairs are almost always visible (Figure 20-23). In long-standing cases, hyperpigmentation is typical. Severe cases can result in scarring and disfigurement from keloid formation (Figure 20-24).

Acne keloidalis nuchae is a similar condition that occurs on the nape of the neck. It also presents with multiple smooth, flesh-colored papules that are quite firm and may coalesce to form keloids (Figure 20-25). Pustules, cysts, and sinus tracts occasionally form. The condition is aggravated by cutting the hair too close to the skin.

**Pathogenesis**

The pathophysiological mechanism responsible for PFB has been clearly elucidated. Essentially, it represents a foreign body reaction\(^{172}\) that is the result of beard hair penetrating the dermis. The tightly curled and undulating surface of hair in black people has several features that facilitate this process. It is elliptical in shape,\(^{173}\) so it develops a pointed tip when shaved. Furthermore, the hair initially grows parallel to the skin surface,\(^{174}\) so that when cut obliquely by shaving, the point is even sharper. Finally, because of its tight curl, the hair has a tendency to grow back toward the epidermis, which is easily penetrated by the pointed hair. The actual penetration of the skin can occur via one of two pathways: through the stratum corneum after growing out from the follicle and arching back toward the skin,\(^{174}\) or by piercing the follicular wall directly.\(^{175}\) The latter results from applying tension to the skin during shaving, thus allowing the sharpened tip to retract under the skin when tension is released.
Fig. 20-25. Acne keloidalis nuchae, recurrent after carbon dioxide laser ablation.

**Diagnosis**

The clinical presentation of PFB is quite characteristic, so the diagnosis is not a difficult one. It is most often confused with acne vulgaris. The absence of comedones (the primary lesions of acne vulgaris), the restriction of disease activity to the beard area, and the history of onset after close shaving all support the diagnosis. Careful visual inspection with magnification will almost always reveal ingrown hairs in some of the perifollicular papules, confirming the diagnosis of PFB.

**Treatment**

Usually, the first phase treatment should be to discontinue shaving and allow beard growth for up to 30 days. By this period of time, most of the ingrown hairs will have been pulled free by the growth of the hair shaft in a direction away from the skin surface.

The next step is to resume shaving. However, the individual should never attempt a close shave: this will facilitate retraction of the sharpened hairs below the skin surface with subsequent penetration of the follicular wall and renewal of the entire process. The simplest mechanism that avoids this problem is shaving with electric clippers, which produces a stubble shave appearance. Excellent results have been reported with this technique,176,177 and it is this author’s treatment of choice. Patients must also be taught to inspect the beard hair closely and to attempt to free any ingrown hairs with a pick. Use of a coarse facial sponge, such as Buf-Puf (manufactured by 3M Products, St. Paul, Minn.), is also helpful for accomplishing this.176,178 At the shaving clinic at Lackland Air Force Base, Texas, this method has been successfully initiated at the time of initial patient presentation, without having to wait the customary 30 days. This allows the patient (usually a basic trainee) to maintain a military appearance and control the problem at the same time.

Shaving can also be accomplished with a manual razor, provided that the proper guidelines are followed (Exhibit 20-6). Careful following of these guidelines is time consuming. With practice, it usually requires about 15 minutes. There are no shortcuts. If an individual is usually rushed in the morning, we often recommend shaving in the evening when he can devote the proper amount of attention that is required.

Successful results have also been reported using electric razors179 and a foil-guarded manual razor specifically designed for patients with PFB (PFB...
Chemical depilatories are also an effective treatment modality. Many patients, however, find them irritating. In addition, they are time consuming and they produce a characteristic odor that many patients find bothersome. A minority of patients seem to prefer this mode of treatment.

Keloids may respond to intralesional steroid therapy, which should be instituted early and aggressively to prevent disfigurement. Severe cases have required excision and grafting. The results of surgical therapy are usually disappointing.

Systemic antibiotic therapy is rarely required in the management of PFB, except in those few cases that become secondarily infected. Some authorities claim efficacy for various topical medications such as tretinoin (applied nightly) and clindamycin (applied twice daily). Occasional use of mild topical steroids may also be useful, particularly for relieving any irritation from depilatories.

Patients must be instructed to avoid plucking of hairs as this may cause breakage of the hair shaft within the follicle with subsequent transfollicular dermal penetration and foreign body response.

Military Considerations

To prevent PFB, one need only grow a beard. Unfortunately, this is contrary to military regulations. In the past, the lack of a consistent policy regarding shaving requirements and the management of PFB from an administrative as well as medical viewpoint turned a relatively minor medical problem into a major social issue. Racial tension over this led to public demonstrations and even frank mutiny by disgruntled black enlisted men in the 1970s. In 1979, one investigator wrote that “pseudofolliculitis barbae has now become the most significant dermatologic disease in the U.S. Army” and some military dermatologists called for “the pan-service acceptance of the voluntary growth of a beard by any service member.”

Fortunately, the lack of a consistent policy was recognized and rectified so that today, PFB clinics are functioning smoothly throughout the military. Administrative guidelines are clearer, commanders are better informed, and medical providers are better educated on the unique problems regarding PFB.

The importance of keeping PFB under control without growing a beard is not simply to maintain a uniform, clean shaven appearance. It is also important for the safety of the individual in a combat environment, where the threat of chemical weapons is ever present. Although tests on the efficacy of gas masks reveal that beard growth of as much as 1/8 in. (up to 3 d) did not significantly alter the seal to the mask, it seems obvious that excessive beard growth would have an adverse effect on the seal. An individual whose PFB was poorly controlled and who required a 30-day profile to allow for beard growth, would likely require evacuation to the rear. Thus, proper management of a common condition like PFB is essential for avoiding unnecessary loss of manpower. The key to the successful management of PFB will always be education and communication among soldiers, healthcare providers, and supervisors.

CUTANEOUS VIRAL INFECTIONS

Herpes Simplex Labialis

Herpes simplex labialis is the condition commonly referred to as “fever blisters.” It is caused by herpes simplex virus (HSV), a double-stranded DNA virus, of which two types exist. Most infections involving the oral mucosa are of type 1, while type 2 is responsible for most HSV infections involving the genitalia (genital herpes is discussed in Chapter 19, Sexually Transmitted Diseases). Like all herpes viruses, HSV-1 is able to persist in a latent form in its host, causing recurrent outbreaks. It occurs worldwide, and by the fourth decade of life more than 90% of the population have developed antibodies to HSV. Thus, recurrent HSV labial infection is an extremely common condition; it affects 20% to 40% of the adult population.

Clinical Features

Pharyngitis and gingivostomatitis are the usual manifestations of primary infection with HSV-1, seen most commonly in children and young adults. Vesicles or erosions may be seen on the palate, gingiva, tongue, lip, or perioral region of the face. Primary infection usually is associated with fever and cervical adenopathy as well, and is most severe in adults.

Recurrent infection manifests as vesicles or erosions on the vermilion border of the lip (Figure 20-26) that usually crust over within 48 hours. Of-
ten, the patient experiences burning and itching at the site prior to vesicle formation. The rate of recurrence varies between 16% and 45%.\textsuperscript{192} Recurrent outbreaks represent reactivation of the virus from its latent form in the trigeminal ganglion.\textsuperscript{192} Although the exact mechanism of reactivation is not known, it is assumed that the virus travels down the axon of the nerve to infect keratinocytes on the adjacent location on the lip. Many precipitating factors have been identified including trauma, UV light exposure, fever, immunosuppression, psychological stress, and exposure to excessive amounts of alcohol, tobacco, or spicy foods.\textsuperscript{193,194}

Recurrent HSV-1 outbreaks can precipitate a reaction known as erythema multiforme, an acute, widespread eruption with characteristic targetlike lesions (Figure 20-27). Antiviral therapy with acyclovir has been shown to prevent relapses of recurrent erythema multiforme.\textsuperscript{195}

\textbf{Diagnosis}

Grouped vesicles or crusted erosions on the lip are strongly suggestive of HSV infection. Lesions that are secondarily infected with bacteria may be difficult to distinguish from primary impetigo, which is characterized by honey-colored crusts. A history of recurrence in the exact location of episodes of minor trauma or stress or UV light exposure support the diagnosis of herpes simplex. Primary gingivostomatitis may resemble aphthae, hand-foot-and-mouth disease (coxsackie virus), Behçet’s syndrome, or Stevens-Johnson syndrome.

The diagnosis of HSV infection is facilitated by performing a Tzanck smear. Material is obtained by scraping the base of a vesicle or erosion. The specimen is then smeared on a glass slide, stained with Wright’s stain, and examined under the microscope. In HSV-infected tissue, multinucleated giant cells and/or keratinocytes with large, pale nuclei with peripheral clumping of chromatin will be seen (Figure 20-28). In addition, culture confirmation of HSV is available through most laboratories. Confirmation of HSV infection through detection of the virus’s DNA by polymerase chain reaction, an exquisitely sensitive technique, has been described\textsuperscript{196} but is expensive and impractical for ordinary use.
Treatment

In most instances, systemic treatment of herpes simplex labialis is not required. Cool compresses and topical application of an antibiotic ointment may promote healing and prevent bacterial superinfection. Topical acyclovir is ineffective in immunocompetent individuals. Patients may benefit from oral acyclovir (200 mg five times daily) if initiated immediately after symptom onset.

Recurrences of herpes simplex labialis can be decreased by prophylactic administration of oral acyclovir (200 mg three times daily). In most cases, the expense of this method of treatment precludes its use. It is indicated, however, for patients with recurrent erythema multiforme due to HSV. It has also been shown to be of practical value in individuals with a history of UV radiation-induced herpes simplex labialis who are going to be exposed to periods of intense UV exposure such as snow skiing (or desert deployment). Zinc oxide, a complete sun blocker, should also be considered for these individuals.

Herpes Zoster

Herpes zoster, also called shingles, is a common neurocutaneous disease caused by the varicella-zoster virus (VZV), a member of the herpesviridae group, which is also the etiologic agent for chickenpox. It occurs in 1.3 to 4.8 persons per 1,000 per year, and is characterized by a vesicular eruption in a dermatomal distribution. Often, it is associated with varying degrees of pain, which may persist beyond the point of healing of the rash.

Clinical Features

Herpes zoster occurs in individuals with a previous history of varicella. Patients typically present

Fig. 20-29. Herpes zoster. (a) Thoracic dermatome. Note sharp cutoff at the midline. (b) Close-up view of (a) to reveal characteristic grouped vesicles on an erythematous base. (c) Lumbar distribution. (d) Sacral distribution with sharp cutoff at midline; note vesicles coalescing and becoming purpuric.
with grouped vesicles on an erythematous base in a dermatomal distribution (Figure 20-29). A generalized eruption may occur in 2% to 5% of patients, particularly in elderly or immunocompromised patients. The eruption is almost always unilateral and rarely extends across the midline. The vesicles may coalesce into larger bullae and may become hemorrhagic or even pustular (Figure 20-30). The patients may or may not experience the prodrome of fever, malaise, and headache. Often, the rash is preceded by radicular pain, a burning sensation, or hyperesthesia in the region of the same dermatome. The most common dermatomes involved are the thoracic, followed by the cranial, cervical, lumbar, and sacral. The most common single nerve involved is the trigeminal nerve—usually a single branch.

New lesions can develop for up to approximately 8 days, but usually do not occur after 4 days from the onset of the rash. Crusting occurs in 10 to 14 days, but lesions may persist for longer periods in elderly or immunocompromised patients. The areas involved may heal with residual postinflammatory hyperpigmentation and can result in significant scarring (Figure 20-31).

**Types of Herpes Zoster**

**Herpes Zoster Ophthalmicus.** Involvement of any branch of the trigeminal nerve is called herpes zoster ophthalmicus (Figure 20-32). Although ocular involvement may occur in association with maxillary or mandibular nerve involvement, this is uncommon. Most frequently, the supraorbital and supratrochlear divisions of the frontal nerve are involved. Hutchinson’s sign consists of lesions on the nasal tip, which indicate involvement of the nasociliary branch of the ophthalmic nerve and a high likelihood of ocular complications. When Hutchinson’s sign is present, immediate ophthalmological consultation is warranted. Eye complications include lid ulceration and scars; conjunc-
Fig. 20-32. Herpes zoster ophthalmicus with marked periorbital and lid edema. Immediate ophthalmological consultation is required.

tivitis; proptosis; keratitis; corneal ulceration, vascularization, or perforation; uveitis; and vascular ischemic injuries.206

Ramsay Hunt’s Syndrome. Ramsay Hunt’s syndrome includes facial nerve palsy as a result of involvement of the geniculate ganglion with VZV (Figure 20-33). Vesicles are usually present on the external ear or tympanic membrane, and auditory symptoms may be present such as tinnitus, vertigo, or diminished auditory acuity. Early intervention with systemic steroids may be beneficial. The facial nerve palsy usually resolves, but can be permanent.

Postherpetic Neuralgia. The acute pain of herpes zoster is usually of a transitory nature. In some individuals, however, particularly the elderly, severe pain may persist for months and become disabling. The frequency of postherpetic neuralgia varies between 15% and 70%.205 It is particularly frequent and severe in patients with herpes zoster ophthalmicus. The neuralgia resolves spontaneously in 50% of patients within 3 months and in 75% within 1 year.208

Diagnosis

The presence of grouped vesicles on an erythematous base in a dermatomal distribution, with associated pain in a similar location, creates little question about the diagnosis of shingles. Disseminated cases may resemble primary varicella infection or disseminated HSV. The Tzanck smear will not differentiate the three. Viral cultures are available, but can take up to 2 weeks to grow. Fluorescein-tagged VZV antibody applied to a Tzanck smear may help to differentiate VSV from HSV but requires a fluorescent microscope.

Treatment

In most cases involving immunocompetent patients, symptomatic treatment will suffice. Cool compresses with Burow’s solution will hasten drying and crusting of the vesicles. For analgesia, narcotics may be required.

Acyclovir has been shown to be effective in shortening the duration of disease and reducing the acute and postzoster pain.209–211 VZV is up to 8-fold less susceptible to acyclovir than HSV,212 so higher doses are required. The recommended dose is 800 mg orally five times daily for 10 days.

The role of systemic corticosteroid therapy in the treatment of VZV infection is somewhat controversial.213 A short course (60 mg/d orally for 5 d tapered over 2 wk) combined with a 10-day course of acyclovir may be useful for the prevention of

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Fig. 20-33. Ramsay Hunt’s syndrome: inability to raise eyebrow with ipsilateral drooping of the corner of the mouth.
postherpetic neuralgia in patients at high risk, such as those greater than 60 years old with VSV involving the trigeminal nerve.

**Molluscum Contagiosum**

Molluscum contagiosum is a benign cutaneous infection caused by a poxvirus. It occurs in children and young adults and is worldwide in distribution. It is spread by fomites as well as by direct contact, including venereal transmission. (Molluscum contagiosum is also discussed in Chapter 19, Sexually Transmitted Diseases.)

**Clinical Features**

The pathognomonic lesion of molluscum contagiosum is a small (2–5 mm), firm, smooth, dome-shaped, translucent papule with a central umbilication. Rapid freezing may accentuate this central depression, thus aiding in the diagnosis (Figure 20-34). With pressure, a white, curdlike substance can be expressed from the center of fully developed lesions. Microscopical examination of a smear of this substance after staining will reveal the characteristic cytoplasmic inclusions known as Henderson-Patterson bodies.

Patients may have few or numerous lesions. They can occur on any cutaneous site, but have a predilection for the pubic region and adjacent locations in adults; in children, the trunk, face, and proximal extremities are most often involved. In general, the lesions are asymptomatic. Without treatment, the disease will last an average of 6 months to 3 years; individual lesions persist for approximately 2 months.

**Diagnosis**

The appearance of typical, small, umbilicated papules on the exposed areas of children or the area of the genitalia of adults usually leaves little doubt as to the diagnosis of molluscum contagiosum. In acquired immunodeficiency syndrome (AIDS) patients, lesions may appear with atypical features. Recently, there have been reports of cutaneous cryptococcosis in AIDS patients resembling molluscum contagiosum, and a case of molluscum contagiosum mimicking a basal cell car-

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**Fig. 20-34.** Molluscum contagiosum. (a) Flesh-colored papules in periorbital location. Note central umbilication of some lesions. (b) Slightly erythematous papules on the trunk. (c) Use of cryotherapy to accentuate the central umbilication, facilitating diagnosis.
cinoma in a patient with AIDS. If any doubt exists, demonstration of the pathognomonic Henderson-Patterson intracytoplasmic inclusions by cytological or histological examination will confirm the diagnosis.

**Treatment**

When approaching the management of molluscum contagiosum, it is important to consider that it is a benign disease in which the lesions eventually resolve spontaneously. Therefore, one should avoid aggressive treatment that may be painful or result in scarring. Some argue that the disease should not be treated at all, particularly in children. Treatment may be justified, however, as an attempt to shorten the course of the disease, thus preventing viral transmission and autoinoculation.

Treatment involves destruction of the lesions by mechanical means such as curettage, cryotherapy, or electrosurgery or by chemical irritants such as retinoic acid, cantharidin, phenol, podophyllin, or trichloroacetic acid. The most effective of these modalities is probably removal by curettage. The treatment may be facilitated in children by use of newly developed topical anesthetic agents such as eutectic mixture of local anesthetic (EMLA).

**Warts**

Warts are benign cutaneous tumors that will affect up to 10% of the population. They are caused by the human papillomavirus (HPV), a virus that has gained a considerable degree of attention due to its ubiquitous nature, tenacity, and oncogenic potential. (Genital warts caused by HPV are discussed in Chapter 19, Sexually Transmitted Diseases.)

The papillomavirus (PV) of the papova group of viruses consists of multiple species that are host-specific such as bovine PV, sheep PV, cottontail rabbit PV (Shope fibroma virus), and HPV. Natural infection by PV is limited to the skin and mucous membranes of the host species. HPV is an ether-resistant virus with icosahedral symmetry and a double-stranded circular DNA. It is the presumed etiologic agent of human warts, based on the facts that DNA and viral particles of HPV have been found in human warts, and filtered extracts of warts will produce papillomas if injected into the skin of human. At least 55 types of HPV have been recognized based on DNA homology, and each type appears to have some degree of anatomic specificity. For instance, HPV type 4 is associated with common warts, while types 6 and 11 are associated with condyloma acuminata of the genital region. Some types apparently have significant oncogenic potential, particularly types 16, 18, and 31, which are commonly identified in cervical carcinoma.

**Clinical Manifestations**

Like all papillomaviruses, HPV can infect the skin and mucous membranes including the genitalia, conjunctiva, oral cavity, and larynx. This discussion is limited to HPV infection of the skin.

**Common Warts.** The common wart (verruca vulgaris) is an exophytic, rough-textured, hyperkeratotic papule that is usually painless. Although typically smaller than 1 cm, warts can form larger plaques. Warts can occur almost anywhere on the skin, but are most common on the dorsum of the hands and fingers, particularly in young children (Figure 20-35). Patients may present with single lesions, but multiple lesions are more common since autoinoculation is a frequent process. Common warts will frequently resolve spontaneously after several months, but they may persist or recur after many years. Warts that resolve spontaneously do not scar. When they occur in the periungual region, they may cause nail plate deformities (Figure 20-36).

**Flat Warts.** Flat warts (verruca plana) are small (1–3 mm), slightly elevated, flesh colored, flat papules that are almost always multiple. They occur mainly on the face, neck, hands, or knees of young adults and adolescents (Figure 20-37), and are resistant to treatment.

**Filiform Warts.** Filiform warts (verruca filiformis) occur most commonly on the face. Multiple lesions may be seen, but they are usually single. These are elongated, thin, pointed projections that, although cornified, remain relatively soft.

**Plantar Warts.** Plantar warts (verruca plantaris) are of two types: mosaic and myrmecia (Figure 20-38). The most commonly seen are the mosaic types, which appear as multiple flat, slightly elevated, hyperkeratotic papules that are usually coalescent. They are most frequently seen in adolescents overlying the metatarsal heads, and are frequently tender and can cause pain on walking. Close inspection of these warts will reveal the typical coarse surface that is sharply demarcated from the surrounding skin. The normal skin markings are interrupted on the surface of the warts. Paring of the keratotic surface will reveal dark, punctate areas that represent thrombosed capillaries. Continued paring will produce pinpoint bleeding.
Fig. 20-35. Common verrucae. (a) Typical grouped warts in a periungual location. (b) Isolated wart on dorsum of hand. (c) Larger wart on thumb; note thrombosed capillaries visible on the surface. (d) Large wart on arm.

Fig. 20-36. Periungual wart causing lifting of distal free edge of the nail plate.
The second type of plantar wart is the myrmecia ("ant hill") type. These are deeper, dome-shaped papules that frequently become inflamed.

**Other Cutaneous Manifestations.** Epidermodysplasia verruciformis is a rare, inherited disease characterized by widespread verruciform plaques that occur in childhood and persist into adulthood. These plaques represent extensive infection with HPV in individuals with impaired immunity. Squamous cell carcinoma will develop in a large number of these patients, particularly on sun-exposed areas.229

The DNA of HPV has been demonstrated in some keratoacanthomas230 and lesions of Bowen’s disease (squamous cell carcinoma in situ).231

**Diagnosis**

The morphology of warts usually is distinctive enough to allow diagnosis by visual inspection. At times they may resemble seborrheic keratoses, actinic keratoses, molluscum contagiosum, cutaneous horns, or acrokeratosis verruciformis of Hopf. Plantar warts may mimic corns (clavi) or calluses. Squeezing a plantar wart usually elicits pain, unlike a callus. In contradistinction to warts, the normal skin markings are accentuated in clavi and calluses. Paring of the warts with a scalpel blade will reveal the characteristic dark, punctate, thrombosed capillaries.

**Treatment**

Like molluscum contagiosum, cutaneous warts are generally a benign infectious process that will frequently resolve spontaneously. Spontaneous remission is seen in 40% of patients within 6 months, and 66% in 2 years.232 Therefore, the physician should avoid overly aggressive treatment that may result in scarring, and be particularly careful to
avoid scarring on the plantar surface as this could cause pain on walking. The modalities available for treating warts are numerous, which speaks for the tenacity of HPV. Recalcitrance is common with warts, and the physician should explain to patients at the initial visit that multiple treatments are sometimes required and recurrences are frequent.

Effective topical keratolytic agents are available. These include salicylic acid, which is available in 40% concentration in a self-adhesive tape (Mediplast, manufactured by Beiersdorf, Norwalk, Conn.) and combined with lactic acid in flexible collodion (Duofilm, manufactured by Stiefel, Coral Gables, Fla.). Both work well under occlusion, if applied daily for several weeks. These may be helpful for treating plantar warts, particularly in children, but can require a fair amount of patience and compliance.

One of the easiest and most effective treatments for warts is curettage with or without electrocoagulation or cryotherapy to the base. Mahrle and Alexander report good results in the treatment of periungual and plantar warts with light focal electrocoagulation, followed by curettage.

Cryotherapy with liquid nitrogen can also be an effective means of eradication. Again, multiple treatments may be required. Bunney et al. report a cure rate of 80% when treating warts on the hands with cryotherapy every 3 weeks for up to 12 weeks.

For warts recalcitrant to the methods already described, particularly periungual or plantar, one might consider intralesional bleomycin or vaporization with the carbon dioxide laser. Facial verruca plana can be most refractory to treatment. Daily application with Retin-A (tretinoin, manufactured by Ortho, Raritan, N.J.) may be helpful, as can twice-daily application of Efudex (5-fluorouracil, manufactured by Roche Laboratories, Nutley, N.J.).

**Military Considerations**

An important consideration for the military physician regarding cutaneous viral infections concerns the role played by stress in triggering outbreaks. The battlefield certainly provides a degree of stress, both emotional and physical, that most people will never experience elsewhere. For this reason, the incidence of infections such as herpes simplex labialis or herpes zoster is likely to be increased. It is extremely important, then, for the military physician to have some useful knowledge regarding their treatment and possible prevention. Some degree of protection may be granted to the individual with a history of UV radiation–induced herpes labialis with the prophylactic use of acyclovir and sun blockers such as zinc oxide. There are no measures available to prevent an episode of shingles. However, the physician can intervene to prevent secondary bacterial infection, which is certainly more likely to occur during wartime, when standards of hygiene are unavoidably lowered. This can be achieved through temporary isolation or hospitalization and the use of topical or systemic antibiotics.

In the treatment of warts, early medical intervention is encouraged to avoid the possible impairment of performance that may occur as the result of widespread involvement of the fingers or the presence of painful plantar lesions. However, overly aggressive treatment should be avoided, because scars or large, iatrogenic bullae could negatively affect performance, permanently or temporarily.

**SUMMARY**

It is imperative for the medical officer to have some basic understanding of the common dermatoses. Historically, these skin diseases have had a substantial impact on the number of casualties suffered during armed conflicts. While usually not considered significant threats to an individual’s health, these maladies will frequently be exacerbated under battlefield conditions, resulting in significant morbidity with a subsequent negative impact on performance of combat duty.

The common dermatoses are usually not difficult to manage. Prompt recognition and intervention by the medical officer may prevent a simple problem from rapidly escalating into an incapacitating condition.

Medical officers with a firm grasp on the recognition and management of common skin diseases will be able to intervene easily to maintain the health of individual troops as well as the collective fighting effectiveness of their units. During armed conflict, given the harsh environmental conditions that exist, intervention may be required frequently. Understanding the information contained in this chapter will enable the medical officer to manage these common dermatoses appropriately and skillfully.
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