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Otezla® (apremilast) is indicated for the treatment of patients with moderate to severe plaque psoriasis who are candidates for phototherapy or systemic therapy.

INDICATIONS

Otezla® (apremilast) is indicated for the treatment of patients with moderate to severe plaque psoriasis who are candidates for phototherapy or systemic therapy.

INDICATIONS AND USAGE

Otezla was evaluated in 2 multicenter, double-blind, placebo-controlled trials of similar design. Patients with moderate to severe plaque psoriasis (N = 297) were randomized 2:1 to Otezla 30 mg or placebo twice daily for 16 weeks, after a 5-day titration.¹

Inclusion criteria: Age ≥18 years, BSA involvement ≥10%, SPGA ≥3, PASI score ≥12, candidates for phototherapy or systemic therapy¹ ¹

PASI-75 response at week 16 (primary endpoint)
- ESTEEM 1: Otezla 33% vs placebo 5% (P < 0.0001)² ³
- Similar PASI-75 response was achieved in ESTEEM 2²

IBSA, body surface area; PASI, Psoriasis Area and Severity Index; SPGA, static Physician Global Assessment.

INDICATIONS

Otezla® (apremilast) is indicated for the treatment of patients with moderate to severe plaque psoriasis who are candidates for phototherapy or systemic therapy.

Otezla is indicated for the treatment of adult patients with active psoriatic arthritis.

IMPORTANT SAFETY INFORMATION

Contraindications

Contraindicated in patients with a known hypersensitivity to apremilast or to any of the excipients in the formulation

Warnings and Precautions

Diarrhea, Nausea and Vomiting: Cases of severe diarrhea, nausea, and vomiting have been reported with the use of Otezla. Most events occurred within the first few weeks of treatment. In some cases patients were hospitalized. Patients 65 years of age or older and patients taking medications that can lead to volume depletion or hypotension may be at a higher risk of complications from severe diarrhea, nausea, or vomiting. Monitor patients who are more susceptible to complications of diarrhea or vomiting: advise patients to contact their healthcare provider. Consider Otezla dose reduction or suspension if patients develop severe diarrhea, nausea, or vomiting

Depression: Treatment with Otezla is associated with an increase in depression. During clinical trials 1.3% (12/920) of patients reported depression, compared to 0.4% (2/506) on placebo. Suicidal behavior was observed in 0.1% (1/1109) of patients on Otezla, compared to 0.2% (1/506) on placebo. Carefully weigh the risks and benefits of treatment with Otezla for patients with a history of depression and/or suicidal thoughts/behavior, or in patients who develop such symptoms while on Otezla. Patients, caregivers, and families should be advised of the need to be alert for the emergence of or worsening of depression, suicidal thoughts or other mood changes, and they should contact their healthcare provider if such changes occur

Weight Decrease: Body weight loss of 5-10% occurred in 12% (96/814) of patients treated with Otezla and in 5% (19/382) of patients treated with placebo. Monitor body weight regularly; evaluate unexplained or clinically significant weight loss, and consider discontinuation of Otezla

Drug Interactions: Apremilast exposure is decreased when Otezla is co-administered with rifampin, a strong CYP450 enzyme inducer; loss of Otezla efficacy may occur. Concomitant use of Otezla with CYP450 enzyme inducers (e.g., rifampin, phenobarbital, carbamazepine, phenytoin) is not recommended

Adverse Reactions

Adverse reactions reported in ≥5% of patients were (Otezla%, placebo%): diarrhea (17, 6), nausea (17, 7), upper respiratory tract infection (9, 6), tension headache (8, 4), and headache (6, 4)

Use in Specific Populations

Pregnancy and Nursing Mothers: Otezla is Pregnancy Category C; it has not been studied in pregnant women. Use during pregnancy only if the potential benefit justifies the potential risk to the fetus. It is not known whether apremilast or its metabolites are present in human milk. Caution should be exercised when Otezla is administered to a nursing woman

Renal Impairment: Otezla dosage should be reduced in patients with severe renal impairment (creatinine clearance less than 30 mL/min); for details, see Dosage and Administration, Section 2, in the Full prescribing Information

Please turn the page for Brief Summary of Full Prescribing Information.


Data includes healthcare professionals (dermatologists, rheumatologists, nurse practitioners, and physician assistants) and their Otezla prescriptions (including refills) from April 2014 through June 2017 for patients with plaque psoriasis or psoriatic arthritis. Source: Data on file, Celgene Corporation.

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Table 3: Adverse Reactions Reported in ≥1% of Patients on OTEZLA and With Greater Frequency Than in Patients on Placebo; up to Day 112 (Week 16)

<table>
<thead>
<tr>
<th>Preferred Term</th>
<th>Placebo (N=506)</th>
<th>OTEZLA 30 mg BID (N=892)</th>
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<tbody>
<tr>
<td></td>
<td>n (%)</td>
<td>n (%)</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>32 (6)</td>
<td>160 (17)</td>
</tr>
<tr>
<td>Nausea</td>
<td>35 (7)</td>
<td>155 (17)</td>
</tr>
<tr>
<td>Upper respiratory tract infection</td>
<td>31 (6)</td>
<td>84 (9)</td>
</tr>
<tr>
<td>Tension headache</td>
<td>21 (4)</td>
<td>75 (8)</td>
</tr>
<tr>
<td>Headache</td>
<td>19 (4)</td>
<td>55 (6)</td>
</tr>
<tr>
<td>Abdominal pain*</td>
<td>11 (2)</td>
<td>39 (4)</td>
</tr>
<tr>
<td>Vomiting</td>
<td>8 (2)</td>
<td>35 (4)</td>
</tr>
<tr>
<td>Fatigue</td>
<td>9 (2)</td>
<td>29 (3)</td>
</tr>
<tr>
<td>Decrease appetite</td>
<td>5 (1)</td>
<td>26 (3)</td>
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<tr>
<td>Insomnia</td>
<td>4 (1)</td>
<td>21 (2)</td>
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<tr>
<td>Back pain</td>
<td>4 (1)</td>
<td>20 (2)</td>
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<tr>
<td>Migraine</td>
<td>5 (1)</td>
<td>19 (2)</td>
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<tr>
<td>Frequent bowel movements</td>
<td>1 (0)</td>
<td>17 (2)</td>
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<tr>
<td>Depression</td>
<td>2 (0)</td>
<td>12 (1)</td>
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<tr>
<td>Bronchitis</td>
<td>2 (0)</td>
<td>12 (1)</td>
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<tr>
<td>Tooth abscess</td>
<td>0 (0)</td>
<td>10 (1)</td>
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<tr>
<td>Folliculitis</td>
<td>0 (0)</td>
<td>9 (1)</td>
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<tr>
<td>Sinus headache</td>
<td>0 (0)</td>
<td>9 (1)</td>
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</table>

*Two subjects treated with OTEZLA experienced serious adverse reaction of abdominal pain.

Severe worsening of psoriasis (rebound) occurred in 0.3% (4/1184) patients following discontinuation of treatment with OTEZLA (apremilast).

DRUG INTERACTIONS

Strong CYP 450 Inducers: Apremilast exposure is decreased when OTEZLA is co-administered with strong CYP450 inducers (such as rifampin) and may result in loss of efficacy [see Warnings and Precautions (5.3) and Clinical Pharmacology (12.3)].

USE IN SPECIFIC POPULATIONS

Pregnancy: Pregnancy Category C: OTEZLA should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus. Pregnancy Exposure Registry: There is a pregnancy exposure registry that monitors pregnancy outcomes in women exposed to OTEZLA during pregnancy. Information about the registry can be obtained by calling 1-877-311-8972.

Nursing Mothers: It is not known whether OTEZLA or its metabolites are present in human milk. Because many drugs are present in human milk, caution should be exercised when OTEZLA is administered to a nursing woman. Pediatric use: The safety and effectiveness of OTEZLA in pediatric patients less than 18 years of age have not been established. Geriatric use: Of the 1257 patients who enrolled in two placebo-controlled psoriasis trials (PSOR 1 and PSOR 2), a total of 108 psoriasis patients were 65 years of age and older, including 9 patients who were 75 years of age and older. No overall differences were observed in the efficacy and safety in elderly patients ≥65 years of age and younger adult patients <65 years of age in the clinical trials. Renal Impairment: Apremilast pharmacokinetics were characterized in subjects with mild, moderate, and severe renal impairment as defined by a creatinine clearance of 60-89, 30-59, and less than 30 mL per minute, respectively, by the Cockcroft-Gault equation. While no dose adjustment is needed in patients with mild or moderate renal impairment, the dose of OTEZLA should be reduced to 30 mg once daily in patients with severe renal impairment [see Dosage and Administration (2.2) and Clinical Pharmacology (12.3)]. Hepatic Impairment: Apremilast pharmacokinetics were characterized in patients with moderate (Child Pugh B) and severe (Child Pugh C) hepatic impairment. No dose adjustment is necessary in these patients.

OVERDOSAGE

In case of overdose, patients should seek immediate medical help. Patients should be managed by symptomatic and supportive care should there be an overdose.

Manufactured for: Celgene Corporation, Summit, NJ 07901

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Pat. http://www.celgene.com/therapies

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**Gauchos to Glaciers!: April 18–27, 2017, Argentina and the Galapagos Islands**

Robert S. Berger, MD

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EDITORIAL

Fake News, Predatory Journals, Case Reports, Biostatistics, and Other Journalistic Musings—Part 1

Warren R. Heymann, MD

We live in an era of information excess delivered by print, digital devices, webcasts, podcasts, and Tweets. Time for reflection is minimal. One of the major precipitants of physician burnout is the endless electronically documenting data, which has replaced speaking to and examining patients, and most importantly thinking about their problems.

Pressure abounds when sorting through the onslaught of data to find what is relevant and true. For those in academia, the need for publication remains undiminished, fueling the cycle further. Although IBM’s Watson can absorb infinite information, unfortunately the glorious human brain can only assimilate a fraction of the data, and its interpretation is compounded by personal bias.

As we have learned, fake news site can generate fabricated stories that can yield devastating consequences when presented as fact.

PREDATORY JOURNALS

Predatory journals have proliferated exponentially, soliciting publications (at a hefty fee). Young academicians may be particularly vulnerable, facing the “publish or perish” scenario. The problem is that many journals will literally publish anything that comes their way. This dilutes the quality of the body of medical literature; indeed, when one is accessing information at point-of-care, the reader can only assume that what was published has been editorially peer reviewed, allowing for at least a modicum of validity.

I receive requests to publish in these journals daily. Although I chuckle when the requests come from other disciplines, such as orthopedics, gastroenterology, or anesthesiology, the situation is not laughable. Out of curiosity, I responded to one journal’s editors, inviting them to visit my website (www.dermatology-insightsandinquiries.com) and choose a contribution that they would find worthwhile. Their response was, “choose whatever you’d like.” Think about that for a moment—whatever I’d like? I can only interpret that as they would publish whatever is sent to them, regardless of the content. How dangerous!

I wholeheartedly concur with the following:

Predatory journals must be deprived of the legitimacy afforded by inclusion in prestigious databases like PubMed. PubMed, operated by the National Library of Medicine, is heavily used by researchers and clinicians to search the medical literature. Efforts by predatory publishers to gain inclusion in the PubMed database through strategies such as purchasing journals that are already PubMed indexed must be thwarted.

Unfortunately, the blog listing predatory journals, started by Jeffrey Beall at the University of Colorado, is no longer available. Cabell International of Beaumont, Texas is planning to develop its own blacklist this year.

IMPACT FACTOR

Journal content is changing, driven by the drive for a higher impact factor. The question is, impact for whom? I read journals for my personal impact factor—how will a paper affect the care I deliver to my patients? I am not decrying the importance of topics such as burden of skin disease or quality of life studies—they are vital for the survival of our specialty as we are facing economic duress. I would just prefer that these
studies be in a new, separate journal devoted to dermatoeco-
nomics.

**CASE REPORTS**

When I read, I go for the clinically relevant studies. I under-
stand why case reports are considered the bottom shelf of the
evidence-based pyramid. Despite that, I get more out of a good
case report and review of the literature than the latest treatise on
the economic burden of hidradenitis suppurativa. Educationally,
I agree that:3–5

Selecting and writing a case report can sharpen critical and
observational skills, improve medical writing, strengthen
the ability to generate and defend a hypothesis, and increase
understanding of patient-centered care. It can give students
valuable experience with the editorial process, motivate them
to take a scholarly view of their clinical work, and allow them
to contribute to the medical literature.3

Should case reports be relegated to electronic publications, they
will not get read unless someone is searching specifically for an
identical case. Increasingly, case reports are published in “open
access” journals for a substantial fee, which may be also burden-
some for the author or their institution.

**CONCLUSIONS**

Dermatologists must become increasingly savvy about how they
acquire new knowledge that affects patient care. Understanding
when medical news is “fake,” appreciating the nature and pitfalls
of predatory journals, considering the “personal” impact factor
of a journal, and reading relevant case reports should enhance
the quality of practice.

Part 2 will discuss biostatistics and other journalistic musings.

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4 Petronic-Rosic V. O Case report: Where art thou?

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COMMENTARY

Media and the Skin

Jacob Johanssen, PhD; Diana Garrisi, PhD

A new study conducted by the University of Texas found that classic Hollywood film villains are statistically more likely to display a skin condition affecting their appearance. The results indicate that six of the all-time top 10 American film antiheroes have dermatologic findings including alopecia (30%), peri-orbital hyperpigmentation (30%), deep rhytides on the face (20%), multiple scars on the face (20%), verrucae vulgaris on the face (20%), and rhinophyma (10%).

Julie Amthor Croley and colleagues dermatologically evaluated the American Film Institute’s 100 Greatest Heroes and Villains list, concluding that skin conditions are used in film to emphasize the dichotomy of good and evil. Their ‘patients’ included the serial killer Hannibal Lecter in The Silence of the Lambs (1991), with androgenic alopecia (Norwood-Hamilton stage 3), and Darth Vader from The Empire Strikes Back (1980), displaying several scars, deep rhytides on the face, peri-orbital hyperpigmentation, and alopecia. The list also included two famous witches—the Wicked Witch of the West (The Wizard of Oz, 1939) and The Queen (Snow White and the Seven Dwarfs, 1937)—both of whom were associated with verruca vulgaris.

Conversely, heroes rarely show conditions severely affecting their appearance. The few exceptions include Rick Blaine (Casablanca, 1942), who has a scar on the lip, and Indiana Jones (Raiders of the Lost Ark, 1981), who has a small chin scar. A notable exception could also be the recent Hollywood superhero Deadpool (2016), who has scars on his face and body.

SKIN AS A STORYTELLING DEVICE

As this research confirms, skin diseases can function to bring the narrative forward in Hollywood storytelling. Bodily deformities are employed not only to visually signify the immorality embodied by the villain, but also to give tangible proof of their evilness. Bodily signs make the antihero iconic and memorable. This is a very simple and quick plot device that aims to give unambiguous information to the audience about who they should identify with. It does not leave space for imagination or interpretation:

the lacerated face of Regan MacNeil in the Exorcist (1973) is not meant to prompt questions; the message is that when you meet the devil, you just know because it shows.

The disrupted face of evil is in direct contrast with the face of the hero or victim, whose immaculacy is exhibited as a reference to their ethical integrity. Skin conditions in cinema are metaphorically presented as a legal proof, the basis upon which the villain must be fought and defeated, leaving the audience happy and comfortable with that choice.

FACIAL DISFIGUREMENT IN THE UK PRESS

Related to the theme of depictions of skin conditions in the media, we are working on a similar project at the Communication and Media Research Institute of the University of Westminster, London. We are currently looking at how facial disfigurement is covered in the UK press. Initial findings show that the news coverage of disfigurement may revert the hero-villain skin dichotomy. How? Most of the stories deals with acid-related attacks.

Acid attacks are a growing problem in the UK: statistics show that the number of admissions to hospital as a consequence of acid attacks rose by 50% between 2005 and 2015.2 The narrative of the newspaper articles builds around the dichotomy of victim-attacker, where the former, whose face shows the wounds and scars left by the corrosive power of acid, is represented as the hero. Scars are then no longer a symbol of moral weaknesses as it happens in movies; on the contrary, the scar becomes the symbol of resistance and bravery. In addition, scars are used to provide documented legal evidence of the physical and psychological pain endured by the person attacked. A wide range of photographs is often displayed in the tabloid press, offering a chronological account of the stages of recovery and surgery from the day of the attack. In this way, the popular press reverses the trope of the face of evil identified by the team of researchers at the University of Texas in cinema. In common, the studies show that mass media still heavily rely on visual binary oppositions to represent human relationships.

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CONCLUSIONS

Reality is seldom based on a binary system as there are gray areas. Even if we assume there is a mutually exclusive distinction between bad and good, it does not necessarily show. This is the reason why organizations such as the UK charity Changing Faces campaign to challenge stereotypical media representations of evilness based on physical appearance, in order to limit the possibilities of increasing the already high rates of prejudice and discrimination surrounding people who have a disfiguring condition. To this end, the UK saw the celebration of Face Equality Day on May 26, 2017. This is the first organized national event created by Changing Faces to raise awareness of facial disfigurement and ensure that everyone is treated fairly and equally, whatever the appearance of their face or body.

REFERENCES


PDs are a heterogeneous group of cutaneous eruptions characterized by pigmented, purpuric macules and papules. Five main clinical subtypes have been described:

- progressive pigmented purpura of Schamberg;
- pigmented purpuric lichenoid dermatosis of Gougerot and Blum;
- purpura annularis telangiectoides of Majocchi (PATM);
- eczematid-like purpura of Doucas and Kapetanakis;
- lichen aureus.1–3

Less commonly described entities include transitory PPD, unilateral linear PPD, granulomatous PPD, and itching purpura of Lowenthal.4,5 Atypical clinical manifestations including unilateral, linear/blaschkoid, zosteriform, and granulomatous variants have also been described.6–8 PPD variants are indistinguishable on histopathology. Schamberg disease is the most common variant, characterized by aggregates of pin-head-sized petechiae resembling cayenne pepper, usually on the lower extremities.9

Multiple triggers have been associated with PPD, including drugs, chemical ingestions, infections, and underlying hematology or systemic diseases. Specifically, multiple different medications have been implicated in the literature as the causative factor responsible for eruptions of PPD. We describe two unique cases of PPD arising in women after the use of an iodine supplement and a vascular endothelial growth factor inhibitor intraocular injection, respectively.

**CASE 1**

A 36-year-old white woman presented with innumerable, min- ute, cayenne-pepper-like pigmented macules diffusely appearing on the trunk and extremities (Figure 1). These asymptomatic lesions had been present for months and were noticed shortly after starting the medication and resolved upon discontinuation. This highlights the importance of considering over-the-counter and intraocular medications when assessing cutaneous eruptions that may be medication related, such as PPDs. (SKINmed. 2018;16:13–17)
CASE 2

An 80-year-old white woman, with a history of Churg-Strauss syndrome, presented with diffuse, nonblanching erythematous macules and petechiae (Figure 4) a few days after the first dose of pegaptanib, a selective vascular endothelial growth factor antagonist, for macular degeneration. A biopsy was taken from a lesion on her arm to confirm the diagnosis. Histopathologic analysis showed attenuation of the epidermis, with basal layer vacuolization, and a patchy lichenoid infiltrate with rare eosinophils and extravasated erythrocytes (Figure 5).

A diagnosis of drug-induced PPD, most likely the pigmented, purpuric lichenoid dermatosis of Gougerot and Blum, was
made. The lesions resolved within a few weeks after completing the pegaptinib intraocular injections.

**DISCUSSION**

PPDs are a heterogeneous group of eruptions with similar histopathologic characteristics. They tend to present clinically with petechiae and golden-brown, pigmented macules, papules, and plaques distributed symmetrically on the lower extremities, but can also be generalized.14 PPD is known to occur in all races, with a predominance in men, except for the PATM variant, which is more common in women.10 Lichen aureus is a rare form of PPD characterized by golden-brown, rust-colored macules or lichenoid papules usually on the lower legs of young adults, especially men, and less frequently in children. This variant may exhibit a segmental pattern, follow the lines of Blaschko or the course of an underlying vein.8

The majority of cases of PPD are benign cutaneous conditions with no systemic involvement, but they can have a chronic and relapsing course. A transitory idiopathic PPD variant resembling Schamberg disease, lasting less than 3 months, has been described in an otherwise healthy young woman, with no history of medication use or systemic disease. These authors noted that the onset of transitory PPD occurred during March–September, and suggested the possibility of an infectious etiology for this variant.11

The PPD variants all share similar histologic features of a perivascular lymphocytic infiltrate, sometimes described as a lymphocytic “capillaritis,” involving superficial vessels in the dermis without frank vasculitis. This perivascular infiltration results in vessel damage and extravasation of erythrocytes into the dermis, with hemosiderin deposition seen in older lesions. A lichenoid infiltrate can be seen in lichen aureus and the Gougerot and Blum variants, and may demonstrate clonal T-cell populations, without other cardinal features suggestive of a frank lymphoma.1,3,12

In a subset of patients, the lesions of PPD progress to cutaneous T-cell lymphoma, most commonly mycosis fungoides (MF), leading some experts to designate PPDs as cutaneous lymphoid dyscrasias.12 Atypical PPDs appear similar to PPDs clinically but histologically show features associated with MF including Sézary cells, laminated dermal sclerosis, and epidermotropism.3,12,13 This leads to diagnostic challenges as there are rare reports of PPD preceding MF, evidence of lymphoid atypia in lesions of typical PPD, and MF with purpuric features described in the literature. A possible existence of PPD/MF overlap has been suggested, with long-standing lesions that look like a reticular form of MF but have the histologic findings of PPD, and these patients require regular follow-up.14

Although most cases of PPD are idiopathic, others have been linked to different factors including drugs, exercise, venous hypertension, gravity, infections, chemical exposures, and systemic illness, such as diabetes and antiphospholipid syndrome.1,15,16 Various medications, food additives, and external contact agents, including clothing dyes, have also been implicated in the etiology of PPD.17–20 Drug-induced cases of PPD are typically transient, lasting up to 4 months, resolve with discontinuation of the medication, and are more likely to involve other body sites. A rechallenge with the same medication would cause a recurrence of the eruptions, facilitating verification of the causative agent in different studies.21,22 A drug-based origin for PPD is therefore determined based on lesional onset and/or resolution after discontinuation of the offending agent. Intraepidermal Sézary cells are less frequent in biopsies of drug-related atypical PPD relative to idiopathic and MF-related atypical PPD, leading to the conclusion that atypical cases of PPD may be of drug-based origin, should not be equated with purpuric MF, and are not necessarily a precursor lesion of MF.12

Common culprit medications include diuretics and nonsteroidal anti-inflammatory drugs, but other drugs have been described including calcium-channel blockers, lipid-lowering agents, β-blockers, angiotensin-converting enzyme inhibitors, antihistamines, antidepressants, analgesics, sedatives, stimulants, antibiotics, cardiovascular drugs, and dietary supplements. Specifically, medications described in association with PPDs include sildenafil, medroxyprogesterone acetate, bufexamac, glipizide, pseudoephedrine, phenothiazine, and creatine supplementation.19,23–26 Most recently, a case of PATM due to isotretinoin has been described.5

Other ingested substances have been implicated in PPD eruptions, as exemplified in a case of widespread PPD lesions induced and rapidly provoked by dietary factors, specifically Coca-Cola and apple cherry fruit spritzer.27 Lichen aureus has been reported in a child after regular consumption of an energy drink, resolving when the drink was stopped.28

Drug-induced PPD has been reported in cases with administration methods including ingestion by the oral route, topical application, and intramuscular and intravenous injections. For example, PPD resulting from the depot contraceptive medroxyprogesterone acetate in a teenage girl completely resolved within 8 weeks of stopping the injections.23 An eruption of PPD due to
glipizide in a male patient resolved within 3 weeks of discontinuing the medication. Bufeaxamac-induced allergic contact dermatitis with hematogenous dissemination presenting with the clinical and histological picture of PPD has also been described. Topical fluorouracil has been associated with a PPD-like eruption. Other medications reported to cause PPD include sedatives such as carbamazepine and meprobamate, vitamin B1, nitroglycerin, chloridiazepoxide, furosemide, and trichlomethiazole.

PPD has been causally linked to pseudoephedrine ingestion. Pseudoephedrine is typically a cause of a fixed drug eruption, especially the nonpigmented variant; however, these authors described a woman with pruritic purpuric plaques with some vesicles after taking 60 mg of pseudoephedrine and 10 mg of loratadine. The patient was rechallenged with loratadine, with negative results. Rechallenge with 15 mg of pseudoephedrine resulted, within 4 hours, in the same eruptions in the same sites as her previous attack. A patch test was also performed using pseudoephedrine and resulted in the same eruption. Biopsies of the positive patch test site and the cutaneous eruption after oral ingestion showed typical findings of PPD. Although PPD eruptions resulting from different administration methods are present in the literature, intraocular medications have not, to the best of our knowledge, been previously described.

Treatment of drug-induced PPD includes stopping the offending medication, which may be sufficient to clear the eruption in most cases. Idiopathic PPD eruptions can be difficult to treat, and the use of topical steroids and compression stockings for the lower extremities in mild cases, with addition of oral vitamin C or pentoxifylline in moderate cases, has been described. In severe or treatment-refractory cases, phototherapy including ultraviolet therapies (PUVA, and narrow-band UVB) and photodynamic therapy has been used, as well as systemic immunosuppressants (PUVA, and narrow-band UVB) and photodynamic therapy. In treatment-refractory cases, phototherapy including ultraviolet therapy after a 1 year of follow-up.

CONCLUSIONS

We report two new causes of PPD, specifically an oral iodine supplement and an intraocular antivascular endothelial growth factor injection. The mechanism of the pathogenesis of drug-induced PPD is not clearly defined, but speculations include a direct toxic effect of the ingested substances or their metabolites on the capillaries, or an immune-mediated vascular injury due to the substance acting as a hapten leading to antigen formation. Other speculations have been made on the effect of medications or their metabolites on the integrity of the vessel wall as part of the pathogenesis of PPD. In drug-induced cases of PPD, stopping the offending medication if possible usually results in prompt resolution of the eruption, as was noted in our patients.

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Botulinum Toxins: Cosmetic and Clinical Applications provides a comprehensive and in-depth review of the use of botulinum toxin for esthetic procedures and medical applications as a stand-alone treatment and as combination therapy with other non-surgical procedures.

- Benefits from a wealth of color images, procedural videos, and expert tips and tricks
- Takes a region oriented approach, providing guidance on treatment of the glabella, forehead, periocular and perioral areas, plus contouring of the lower face, lower leg and calf, and neck rejuvenation
- Contains a thorough review of non-cosmetic treatments such as correction of facial asymmetry and treatment of axillary hyperhidrosis, plus palm, sole, and craniofacial hyperhidrosis
- Covers exciting new topics, such as future injectables, topical botulinum toxin, and facial contouring including treatment for benign masseter hypertrophy
- Includes considerations for darker skin types

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ORIGINAL CONTRIBUTION

Salivary Cortisol and Salivary Flow Rate in Clinical Types of Oral Lichen Planus

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ABSTRACT

Patients with oral lichen planus (OLP) may develop immune sialadenitis that causes a dry mouth. The role of cortisol in autoimmune diseases is well known; yet studies on this subject are controversial. In this study, the salivary flow rate and salivary cortisol level were compared among subtypes of OLP. This study involved three groups of patients: (1) 11 with reticular OLP, (2) 20 with atrophic-erosive OLP, and (3) 30 with no apparent oral lesion. The salivary flow rate in the control group was significantly higher than in OLP patients. The mean level of cortisol in atrophic-erosive cases was higher than in reticular cases and in the control group; however, there was no significant difference between the three study groups. In a previous study, there was moderate to severe acinar atrophy in two-thirds of patients with OLP, which may explain the decreased salivary flow rate in these patients. (SKINmed. 2018;16:19–22)

Studies have evaluated the salivary cortisol level in individuals with lichen planus, but there is no comprehensive study evaluating the salivary cortisol level in subtypes of OLP. Higher autoantibody levels in autoimmune connective tissue diseases are accompanied by elevated cortisol levels. It can be concluded that, in the atrophic-erosive type of OLP, higher salivary cortisol concentrations may reflect higher circulating autoantibody levels. The flow rate of saliva was lower in OLP subjects compared to control participants, which suggests hyposalivation in OLP patients. Although the salivary cortisol level was relatively higher in patients with OLP, a significant association between salivary cortisol level and OLP was not observed.

Lichen planus is a chronic inflammatory mucocutaneous disease, which frequently manifests in the oral mucosa. Oral lesions are categorized as reticular, papular, bullous, erosive-ulcerative, atrophic (erythematous), and plaque-like lesions. Xerostomia and decreased salivary flow rate have been reported in OLP, but controversies exist regarding this. OLP patients may develop immune sialadenitis that causes a dry mouth, which was at first thought to be Sjögren’s syndrome. Therefore salivary gland dysfunction in OLP has been proposed. One of the aims of this study was the evaluation of salivary flow rate in such patients.

Xerostomia may exist despite normal salivary flow rate. Changes in the quality of saliva that occur in salivary gland dysfunction can cause xerostomia. One of the possible salivary composition-al changes may be of cortisol; cortisol plays a significant role in regulating the immunologic and inflammatory responses. Salivary cortisol level is a better indicator than the serum cortisol level because salivary cortisol is not bound to albumin or corticosteroid-binding globulin, as it is in the serum. In addition, collection of saliva samples is more convenient and noninvasive.

Although the role of cortisol in autoimmune diseases is well known, studies on this subject have been controversial. Previous studies have assessed the salivary level of cortisol in lichen planus, but there is no study evaluating the salivary level of cortisol in different subtypes of this disease. Therefore this study compared salivary cortisol level among atrophic-erosive and reticular types of OLP.

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MATERIALS AND METHODS
This case-control study involved 61 patients seen in the department of oral and maxillofacial medicine. All participants provided written informed consent before samples were collected. The participants were divided into three groups:

- Group 1: 11 patients with histopathologically confirmed reticular-type OLP, with a mean age of 54 years.
- Group 2: 20 patients with histopathologically confirmed atrophic-erosive OLP, with a mean age of 51 years.
- Group 3: 30 patients with no apparent lesion on the oral mucosa, with a mean age of 40 years.

Patients with systemic diseases, autoimmune disorders, hepatitis, HIV, infection, renal disease, oral lichenoid reactions, or a history of taking autoimmune, systemic, or local medications were excluded from the study, as were pregnant women and nursing mothers.

Unstimulated saliva samples were collected between 9 and 11 AM. All the subjects were asked to refrain from oral intake, tooth brushing, smoking, and using any mouthwash for at least 90 minutes before collection of the saliva sample. During sample collection, the lips were cleaned of any cosmetic products. Participants were first asked to stay in a resting position and swallow all the saliva in their mouth, lean their heads forward for 15 minutes, and then spit the accumulated saliva into the sterilized tubes. The samples were then centrifuged at 2000 rpm for 10 minutes. The supernatant fluid was frozen at −20°C until further analysis. The salivary cortisol level of the samples was determined using a salivary cortisol enzyme-linked immunosorbent assay kit.

Quantitative data were presented as means±SD. One-sided variant analysis was used for the comparison between the three study groups. The Games-Howell test was used for paired comparison between study groups. The level of significance was set at <.05.

RESULTS
The basic characteristics and the mean of unstimulated salivary cortisol level and salivary flow rates in the study groups are displayed in the Table. The means and 95% confidence intervals for unstimulated salivary cortisol level and salivary flow rates are shown in Figures 1 and 2.

Although the mean level of cortisol in atrophic-erosive disease was higher than in reticular disease and in the control group, there was no significant difference in the mean cortisol level between the three study groups (P=.39). In addition, the flow of unstimulated saliva was not the same in the study groups (P=.002), in that the flow rate in the control group was significantly higher than that in OLP patients. There was no significant difference between the two subtypes of OLP (groups 2 and 3) regarding salivary flow rate (P=0.87).
DISCUSSION

Reticular is the most common subtype of OLP; however, more patients with atrophic-erosive than reticular disease were included in this study. This might be because the pain and burning sensation seen in the atrophic-erosive disease lead to more patients being referred to dental clinics.

The salivary flow rate in OLP patients was significantly less than that in controls, but there were no significant differences between the two subtypes of OLP. The different findings in the salivary flow rate of OLP in other studies might be explained by differences in saliva collection methods and times of collection. It has been determined that unstimulated salivary flow rate is regulated by a circadian rhythm, being low at approximately 6 AM and high at around 6 PM. Therefore, in the current study, the time of collection was set to be the same for all participants (between 9 and 11 AM).

In the previous study, labial salivary gland biopsy showed moderate to severe acinar atrophy in two-thirds of patients with OLP, which is consistent with our result; however, sample size was small, and further studies are needed to find a relationship between structural changes in salivary glands and diminished salivary flow rate in OLP.

Oral mucosa and skin have the same wound healing phases, but healing occurs more rapidly and with less scar formation in the oral cavity. Several factors influence the faster wound healing of the oral mucosa. For example, there is a higher baseline turnover of cells in the oral cavity than the skin. The presence of saliva is another important factor. Saliva creates a humid environment and contains important proteins that play a role in epithelial healing, such as epidermal growth factor, secretory leukocyte protease inhibitor, and especially histatin. Thus a decrease in salivary flow rate may affect the healing of injured or lesional mucosa, which can exacerbate the chronicity of OLP. An assessment of hyposalivation in OLP may therefore be beneficial for treatment as well as understanding of disease etiology.

The cell-mediated immune system provides the most important contribution to the development of OLP. Cortisol plays a significant role in regulating the immune system response and activating the sympathetic nervous system. Salivary cortisol tracks the circadian variation of serum cortisol, with the highest levels in the morning.

In the current study, salivary cortisol level was higher in patients with OLP than control participants. In addition, salivary cortisol level was higher in atrophic-erosive disease than reticular disease, although there was no significant difference between the three study groups (P > .05). The higher salivary cortisol level in the OLP group was in accordance with the results of other studies.

In one study, salivary cortisol levels were also higher in OLP but did not differ significantly between the OLP group and controls. It seems that the finding of an insignificant or significant difference between OLP and control groups is related to the participants’ psychological state. For example, in another study, cortisol levels were significantly higher in the OLP group, and these patients had significantly higher depression, anxiety, and stress scores.

Higher autoantibody levels in autoimmune connective tissue diseases such as systemic lupus erythematosus are accompanied by elevated cortisol concentrations. It can be concluded that the higher salivary cortisol levels of atrophic-erosive OLP may reflect higher circulating autoantibody levels. A study of 320 OLP patients showed that erosive OLP was the risk factor contributing to the detection of serum antinuclear antibody and anti-thyroglobulin antibody anti-thyroid microsomal antibodies in patients with OLP.
The present study has some limitations, such as a lack of serum cortisol measurements. It is recommended that more extensive study groups of OLP and control cases should be evaluated, with the same number of female and male participants within each group.

CONCLUSIONS

Although the salivary cortisol level was relatively higher in patients with OLP, a significant association between salivary cortisol level and OLP was not observed. The flow rate of saliva was significantly lower in OLP. Further studies of structural changes of salivary gland in OLP are recommended.

REFERENCES

The population for this study included 52% boys and 48% girls, the median age being 7 years old, with the youngest patient being 2 months old and oldest 17.9 years old. The most common hospitalizations included scabies (32%), atopic dermatitis (AD; 31%), dermatitis, not specified (4.6%), dermatitis herpetiformis (3.9%), scleroderma (4.25%), and alopecia areata (2.48%). In 83% of cases, the children were hospitalized only once. The most common cause of recurrent admissions was AD; however, one patient presented with six admissions due to erythroderma, and one patient had five admissions due to mycosis fungoides. AD and scabies are the most common causes of admission to the dermatology department, and are a major concern for the pediatric dermatology population in the Pomeranian region of Poland.

Pediatric dermatology is a narrow field of medicine, often presenting a great challenge to dermatologists as well as pediatricians in the approach to treating patients. Up to 30% of pediatric primary care visits involve skin-related diseases. It is common practice for children presenting with a skin eruption or pruritis to be seen initially by a pediatrician rather than a dermatologist. The importance of the dermatologic manifestations should, therefore, be stressed to pediatricians, as should a good level of communication across departments.

With growing body image pressure on teenagers from the media, skin conditions play a key role in self-worth and normal development. Several centers report epidemiologic data for pediatric dermatologic manifestations. The geographic differences most often seen indicate the importance of localization and management. One study has reported AD, with an incidence of 25.9%, as the most common in its epidemiologic study, followed by pigmented nevi (9.1%) and verrucae (5.0%) for outpatient admissions. In another study, AD was the most common disease, with a frequency of 59.5%, followed by 7.1% seborrhoeic dermatitis, and 4.2% superficial mycoses. The Department of Dermatology, Venereology and Allergology of the Medical University of Gdansk is a specialized center admitting patients of all ages from the entire Pomeranian region. Patients who present with difficult diagnostic and therapeutic challenges are often referred by local pediatricians or other hospital departments.

**MATERIALS AND METHODS**

We retrospectively analyzed 282 pediatric patients admitted to the Department of Dermatology and Allergology of the Medical University of Gdansk over the 3-year period from January 2013 through December 2015.
Personal identification number and initial diagnosis were collected by department nurses upon admission to the hospital and recorded in the admission book. Collected data were further compared with the patient’s electronic records from the hospital electronic medical database—CliniNet. The age of the patients was deduced from the personal identification numbers. The final diagnosis determining our study was obtained from the CliniNet medical record database. The statistical data included patients aged from 2 months to 17 years 9 months, with a median age of 7 years old. We recorded the personal identification number, number of hospitalizations, dates of hospitalizations, age at time of admission, final diagnosis, age of initial diagnosis, other associated conditions/past medical history, family history of atopy, mode of delivery, and APGAR score.

All data were obtained according to the Helsinki Declaration of 1975; the analysis was carried out ethically. Standard diagnostic criteria were used, and rare cases were also recorded, all in a confidential manner. Programs used to store and present the data were Numbers version 4.0.5 and GraphPad Prism 7.

**Figure 1.** Prevalence of the most common pediatric dermatologic hospital admissions over the 3-year study period, 2013 to 2015.

**Figure 2.** Frequency of multiple pediatric dermatologic hospital admissions accordingly to etiology over the 3-year study period, 2013 to 2015. Abbreviation: AD, atopic dermatitis.
RESULTS

Of the 282 participants, 52% were boys and 48% were girls. The median age was 7 years old, the youngest being 2 months old and oldest 17 years 9 months old. The most common hospital admissions included scabies (32%; mean age 7.24 years), AD (31%; mean age 7.0 years), psoriasis (12%), dermatitis, not specified (4.6%), dermatitis herpetiformis (3.9%), scleroderma (4.2%), alopecia areata (2.48%), and vasculitis (0.98%) (Figure 1). In 83% of cases, the children were hospitalized only on one occasion.

The importance of the causes of multiple hospitalizations was also noted (Figure 2). Despite scabies being the most common pediatric condition for readmission, readmission for AD is of more concern, with an average two visits to our department, implying the persistence of the disease and difficulty in treatment. We have also reported one patient with erythroderma which required six hospital admissions, and one patient with mycosis fungoides, admitted five times.

On further analysis of the yearly disease frequency, we found that AD predominated in 2013 and 2015, contrary to 2014, where an increase in scabies was observed. The number of psoriasis admissions increased consistently over the years, and we interestingly also discovered for 2015 a surprisingly higher number of cases of scleroderma, a condition that is exceptionally rare among the pediatric population.

Upon detailed investigation of the population with scabies, among the 91 subjects, 10 had associated AD in their past medical history records, 15 were also diagnosed with AD by our department at the time of hospitalization, and had patients had associated asthma. Asthma, chronic rhinitis, and AD fall into the atopy category—the tendency to hyperallergy and the production of immunoglobulin E due to genetic predisposition, resulting in 33% of cases of scabies being associated with atopy.

Of the patients with AD, 31% had two associated skin conditions—dermatitis herpetiformis and psoriasis—as the most frequent co-morbidities. Finally, in up to 7.9% of children admitted to the department with other skin manifestations, AD was also noted in their past medical history Other diseases accompanying AD in the final diagnosis included predominantly asthma-associated upper respiratory tract infection in 15.7%.

DISCUSSION

Establishing the right diagnosis is crucial to prevent adverse drug reactions and unfavorable clinical outcomes during a course of immunosuppressive treatment for scabies infections. Many patients in our clinical are hospitalized as a result of misdiagnosed non-life-threatening disease and inappropriately applied treatment regimens.

The prevalence of childhood AD in the general European population approaches up to 20% in elementary school children. The disruption of the antimicrobial barrier creates an ideal environment for the staphylococcal infections. A hygiene hypothesis has been proposed, and a negative correlation between family size and AD has been revealed, in contrast to scabies infections in overcrowded settings such as nurseries and hospital wards. One study highlights the importance of “crusted” scabies—the hyperproliferation of the mites causing scabies can lead to hyperkeratosis of the skin, which often is localized and evident in an immunocompromised patient—ie, after prolonged corticosteroid usage. Several factors to consider are misdiagnosis, improper corticosteroid use, education on steroid stepdown protocols or inadequate patient education on a particular skin condition, and poor compliance with treatment.

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Any kind of hypotrichosis or atrichia developing in a young couple’s infant can be a major source of psychological stress for the family as a whole. For the dermatologist, it is important to be aware of the wide variety of conditions that can be associated with congenital hypotrichosis or atrichia. There have been many different attempts to classify congenital hypotrichosis. One of the most accepted and practical follows the classification of congenital alopecia and hypotrichosis proposed by Camacho.

The Camacho classification broadly divides congenital alopecia and hypotrichosis into generalized and localized hypotrichosis. Recent developments in the field of genetics, especially whole-exome and whole-genome sequencing, have led to a better understanding of genetic hypotrichosis.

**TERMINOLOGY**

Unlike alopecia, which describes hair loss in a previously hairy area, the term congenital hypotrichosis refers to the absence of hair growth in the first place. Most described cases of hypotrichoses are associated with genetic defects, many of which have now been pinpointed. Congenital atrichia (also called papular atrichia) is associated with the presence of normal hair at birth, which is gradually lost during the course of childhood and never regrows. Congenital atrichia is also considered to be essentially a genetic defect, which leads to a breakdown in communication between the dermal papilla and the hair.

**CONGENITAL ATRICHIJA**

Atrichia congenita with papular lesions (APL) is a group of genetic conditions characterized clinically by complete and irreversible hair loss shortly after birth. APL is associated with the development of keratin-filled cysts, giving the papular appearance. The disease has been reported in Indian, Irish, and Arab Israeli populations. Inheritance is usually autosomal recessive, although sporadic, dominant, and irregular dominant inheritance have been reported.

The exact etiology is not known, but homozygous mutations in the ‘hairless’ gene (HR) have been suggested to be one of the possible factors involved. HR genes are located on chromosome 8p21.2 and encode a putative single zinc finger transcription factor protein, which is important in regulating the normal hair cycle. The mutation diminishes DNA-binding activity, disturbing hair cycle events. The hair matrix cells in APL undergo premature apoptosis and separation from the epithelial sheath. Communication between the dermal papillae and stem cells in the bulge are not transmitted, leading to cessation of hair growth.

It is important to differentiate conditions like APL from early-onset alopecia universalis, and useful diagnostic criteria for APL have been proposed. These include five major criteria (of which four are required for diagnosis) and five supplementary criteria. The major criteria are: permanent and complete absence of scalp hair by the first few months of life; few to widespread, smooth, whitish, or milia-like papules on the face, scalp, arms, elbows, thighs, or knees from infancy or childhood; replacement of mature follicle structures by follicular cysts filled with cornified material on scalp histology; mutations in the HR gene; and clinical and or molecular exclusion of vitamin D–dependent rickets (as mutations in the vitamin D receptor gene associated with rickets have been found to produce a phenotypical picture similar to

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**ABSTRACT**

A wide range of conditions can present with congenital hypotrichoses/atrichia. Awareness of these conditions can help in the proper and timely diagnosis and counseling of affected families, and in some cases avoid unnecessary investigations. The rapid growth in genetic analysis of diseases has also led to an increased knowledge of the genetic and molecular basis of many of these conditions. This contribution briefly reviews updates on some of the most common conditions associated with congenital hypotrichosis/atrichia. (SKINmed. 2018;16:27–32)
the atrichia associated with \textit{HR} gene mutations\textsuperscript{a}). The minor criteria are: family history of consanguinity; absence of secondary axillary, pubic, or body hair growth and/or sparse eyebrows and eyelashes; normal growth and development, including normal bones, teeth, nails, and sweating; whitish hypopigmented streaks on the scalp; and lack of response to any treatment modality.\textsuperscript{4}

Dermatoscopy of the scalp (trichoscopy) might be another useful tool for diagnosing APL and differentiating it from alopecia universalis. Dermatoscopy of completely developed APL will show absent follicular orifices, without features such as black/yellow dots and exclamation mark hairs, which can be seen in variants of alopecia areata.\textsuperscript{3}

**CONGENITAL HYPOTRICHOSIS**

**Generalized types**

The Table summarizes types of congenital hypotrichosis.

**Hereditary hypotrichosis simplex of the scalp**

This rare autosomal dominant genotrichosis is caused by mutations in the \textit{CDSN} gene encoding corneodesmosin. \textit{CDSN} codes for a keratinocyte adhesion molecule expressed in the inner root sheaths of hair follicles and has been found to be physiologically important for hair follicle function. The mutation causes expression of an abnormal protein that might be toxic to hair growth. The disease has been reported in Chinese, Indian, and Mexican families.

The hairs are normal in the first years of life, after which patients develop progressive, gradual loss of scalp hair that starts in the middle of the first decade of life and progresses to almost complete baldness by the third decade. Body hairs are spared. Occasionally, patients complain of scalp pruritus. The disease has been reported in consanguineous families.

**Hereditary hypotrichosis simplex**

This is a rare autosomal dominant form of hair loss characterized by hair follicle miniaturization. It is caused by mutations in the \textit{APCDD1} gene, encoding adenomatous polyposis coli down-regulated 1, which is expressed in the epidermal and dermal compartments of the hair follicle, and has a role in hair follicle miniaturization. \textit{APCDD} is expressed in the human scalp. The disease has been reported in Italian, Chinese, and Pakistani families.

Hairs are normal or sparse at birth and then start to thin and grow only slowly, leaving the patient with short body and scalp hair. Eyelashes, eyebrows, and male beards are usually spared. Occasionally, hairs are light-colored with a hypopigmented hair shaft and a tapered end. There are no other systemic anomalies. Under light microscopy, the bulb portion of the plucked hairs may show dystrophic features and evidence of miniaturization.\textsuperscript{14,15}

**Autosomal recessive hypotrichosis with wooly hair**

This autosomal recessive disorder is described in consanguineous Pakistani families. The disease is caused by mutations in either the \textit{LIPH} gene, encoding lipase H, or the \textit{LPAR6} gene, encoding lysophosphatidic acid receptor 6; these are expressed in the inner root sheath of the hair follicle and are important in hair follicle growth and differentiation. Patients present with hypotrichosis or complete loss of scalp hair, along with wooly hair. Eyelashes and body hairs are reported to be sparse in some cases, and there sparing of the beard hairs in males.\textsuperscript{16,17}

**Localized autosomal recessive hypotrichosis or autosomal recessive hypotrichosis simplex**

With this group of autosomal recessive disorders, patients have fragile scalp, sometimes trunk, and extremity hairs that break easily, leaving sparse, short hairs. Facial, axillary, and pubic hair are usually spared. Occasionally, patients complain of scalp pruritus.

These hair changes are described with three different mutations. Desmoglein 4 is a member of the desmosomal cadherin family that is expressed in the hair follicle and suprabasal epidermis; it is important in hair follicle proliferation and differentiation. Mutation of the \textit{DSG4} gene has been reported in a group of Pakistani and Iraqi/Iranian families.\textsuperscript{18–20} Mutation in either the \textit{LIPH} or the \textit{LPAR6} gene can cause similar features in the absence of wooly hair (as the mutation is at a different locus). Electron and light microscopic features are variable between patients, and include thick and thin hair parts on light microscopy, and longitudinal grooving with absent cuticular cells on electron microscopy.\textsuperscript{20,21}

**Marie Unna hereditary hypotrichosis**

This is an autosomal dominant disorder caused by a mutation in the untranslated region of the \textit{HR} gene. This region is an upstream regulator of the \textit{HR} gene located on chromosome 8p21.\textsuperscript{22–24} The disease has been described in Europe, North America, India, and China.

Hypotrichosis is distinctive at each stage of life of affected individuals. At birth, affected individuals have absent hair followed by regrowth of coarse, twisted, wiry hair (a diagnostic clue) in childhood, and then progressive nonscarring loss of hair on pu-
<table>
<thead>
<tr>
<th>Syndrome</th>
<th>Mode of Inheritance</th>
<th>Gene Involved</th>
<th>Hair Abnormalities</th>
<th>Associated Features</th>
<th>Histopathology</th>
<th>Light Microscopy</th>
<th>Electron Microscopy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hereditary hypotrichosis simplex of the scalp</td>
<td>AD</td>
<td>CDSN</td>
<td>Hairs normal in the first years of life. Progressive loss of scalp hair starting in the middle of the first decade of life. Almost complete loss of scalp hair by the third decade. Body hairs spared</td>
<td>None</td>
<td>Decreased number of hair follicles with faint chronic, predominantly lymphocytic perifollicular infiltration or no inflammation</td>
<td>Normal</td>
<td></td>
</tr>
<tr>
<td>Hereditary hypotrichosis simplex</td>
<td>AD</td>
<td>APCDD1</td>
<td>Normal or sparse scalp hairs at birth. Thinning and slow growth leaving the patient with short scalp and body hairs. Eyelashes, eyebrows, and male beards are usually spared. Light-colored hairs and tapered shafts</td>
<td>None</td>
<td>Dystrophic features and evidence of miniaturization involving hair bulb</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Autosomal recessive hypotrichosis with wooly hair</td>
<td>AR</td>
<td>LIPH or LPAR6</td>
<td>Hypotrichosis or complete loss of scalp hair with wooly hair. Eyelashes and body hairs sparse in some cases, with sparing of beard hairs in males</td>
<td>None</td>
<td></td>
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</tr>
<tr>
<td>Localized autosomal recessive hypotrichosis or autosomal recessive hypotrichosis simplex</td>
<td>AR</td>
<td>DSG4 LIPH LPAR6</td>
<td>Fragile scalp hairs, sometimes trunk and extremity hairs. Sparse short hairs. Facial, axillary, and pubic hair usually spared. Scalp pruritus</td>
<td>None</td>
<td>Thick and thin hair parts. Longitudinal grooving with absent cuticular cells</td>
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</tr>
<tr>
<td>Marie-Unna hereditary hypotrichosis</td>
<td>AD</td>
<td>Untranslated region of HR</td>
<td>Absent hair at birth followed by regrowth of coarse, twisted, wiry hair, followed by progressive nonscarring loss on puberty. Body hairs including eyebrows and eyelashes are sparse or absent</td>
<td>Rare Milia-like facial lesions, tooth abnormalities, limb abnormalities, Ehlers-Danlos syndrome, and juvenile macular degeneration</td>
<td>Mild to moderate inflammatory infiltrate with reduced number of hair follicles and lack of scarring. Hair shafts deeply pigmented with variations in hair shaft diameter, and irregularly twisted hairs.</td>
<td>Peeling of the cuticle occasionally seen</td>
<td></td>
</tr>
<tr>
<td>Hypotrichosis with juvenile macular dystrophy</td>
<td>AR</td>
<td>CDH3</td>
<td>Short and sparse scalp hair. Progressive macular dystrophy</td>
<td>Increased catagen-telogen hair follicles. Increased ratio of vellus hair follicles to terminal hair follicles. Lack of inflammation or scarring</td>
<td></td>
<td></td>
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</tbody>
</table>

Continued


**Mari hypotrichosis**

This autosomal recessive disorder is described in the Mari population, a Finno-Ugric ethnic group who have traditionally lived along the Volga and Kama rivers of Russia. Patients present with hypotrichosis of scalp hair after birth, and any hairs left are usually wiry and twisted. Eyebrows and eyelashes are gradually lost within the first year of life. Pubic and axillary hairs are absent, and other body hairs are thin and sparse. Rarely, patients present with follicular hyperkertosis.28,29

The disease is attributed to a mutation in the LIPH gene, expressed in hair follicles. This gene encodes membrane-associated phosphatidic acid-selective phospholipase A1α, which is important in regulating the production of bioactive lipids important in hair growth and development.30

**Hypotrichosis with juvenile macular dystrophy**

This autosomal recessive disorder is described in Muslim Arab Israeli populations and some Portuguese families. It is caused by loss-of-function mutations in CDH3, which encodes P-cadherin, a component of the adherent junction in the retinal pigment epithelium and hair follicles; it is also important in regulating cell signaling, cycling, and growth in the hair follicles. Hair abnormalities include short and sparse scalp hair along with progressive macular dystrophy leading to blindness. Increased catagen-telogen hair follicles, an increased ratio of vellus hair follicles to terminal hair follicles, and lack of inflammation or scarring may be observed on histopathology.31–33 Ectodermal dysplasia, ectrodactyly, and macular dystrophy syndrome are caused by the same mutation.34

**Cartilage-hair hypoplasia**

This is an autosomal recessive disorder caused by a mutation in the RMRP gene, encoding RNA component of mitochondrial
RNA processing endoribonuclease, which is involved in ribosomal assembly and cell cycle regulation. The disease is common in the Amish community in the United States. It has been also reported in Indian, Finnish, and Japanese populations.

Patients usually have short stature and short limbs, along with joint laxity and mobility issues. In most cases, affected patients have immunodeficiency and increased risk of malignancy (non-Hodgkin lymphoma, basal cell carcinoma). Other associated clinical manifestations are anemia, defective spermatogenesis, and gastrointestinal abnormalities. The scalp shows sparse thin hairs with blonde eyebrows and eyelashes. Hair shafts lack a central pigment core. Other conditions associated with localized hypotrichosis include nevoid conditions such as nevus sebaceous.

**CONCLUSIONS**

Congenital atrichia and hypotrichia are relatively rare conditions. APL is the most common presentation of congenital atrichia, while congenital hypotrichia includes a group of conditions, broadly classified into localized and generalized types. Advances in genetic studies have led to an improved understanding of the pathogenesis of conditions characterized by congenital hypotrichosis and congenital atrichia. Further research is warranted to explore treatment options based on the molecular basis of these conditions.

**REFERENCES**


SELF ASSESSMENT EXAMINATION

W. Clark Lambert, MD, PhD

Instructions for questions 1–5: For each numbered question, choose the one best numbered response, unless directed otherwise.

1. Congenital hypotrichosis is distinguished from alopecia by:
   a. Hair loss in a previously hairy area.
   b. Absence of hair growth in the first place.
   c. Dense, curly hair instead of normal hair.
   d. Any of the above.
   e. None of these.

2. Alopecia is distinguished from congenital hypotrichosis by:
   a. Hair loss in a previously hairy area.
   b. Absence of hair growth in the first place.
   c. Dense, curly hair instead of normal hair.
   d. Any of the above.
   e. None of these.

3. Most described cases of hypotrichosis are associated with:
   a. Genetic defect(s).
   b. Metabolic defect(s) due to malnutrition.
   c. Metabolic defect(s) due to zinc deficiency.
   d. Metabolic defect(s) due to radiation in the environment.
   e. Metabolic defect(s) due to unknown causes.

4. Disorders sometimes associated with one or more of the various types of generalized hypotrichosis or atrichia include:
   a. Ehlers-Danlos syndrome.
   b. Juvenile macular degeneration of the eye.
   c. Limb abnormalities.
   d. Milia-like facial lesions.
   e. Tooth abnormalities.
   f. All of the above.

5. Which one of the following is most prone to mimic congenital atrichia (atrichia congenita with papular lesions, or APL)?
   a. Vitamin A deficiency.
   b. Vitamin C deficiency.
   c. Vitamin D deficiency.
   d. Vitamin E deficiency.
   e. Vitamin K deficiency.

ANSWERS TO EXAMINATION:
1. b; 2. a; 3. a; 4. f; 5. e.
Edward L. Keyes Resident Contest for Outstanding Case Reports

13th World Congress of the International Academy of Cosmetic Dermatology
Dubrovnik, Croatia June 28–July 1, 2018

Abstract deadline: March 31, 2018

To be awarded for the best Case Report submitted by a physician in training (resident, fellow, or registrar) for presentation at the 13th World Congress of the International Academy of Cosmetic Dermatology in Dubrovnik, Croatia from June 28–July 1, 2018.

We invite you to submit original Case Reports that reflect the presentation of new ideas and original observations to the Academy membership and other attendees of the Congress. The case may be medical, surgical, and cosmetic (or combined) in nature.

The author whose abstract obtains the highest score during the review process will receive a scholarship by the IACD to present the full paper at the 13th World Congress of the International Academy of Cosmetic Dermatology in Dubrovnik, Croatia from June 28–July 1, 2018. The scholarship will provide reasonable travel expenses, lodging for 3 nights, the Congress registration fee, and a basic spending stipend.

Please submit your case report abstract via email to vrosic@medicine.bsd.uchicago.edu before noon, CDT, March 31, 2018. The abstract should be no longer than 2,500 characters including spacing. Material that was previously presented, published, or submitted for publication should not be offered. Applications will be graded based upon the educational value of the abstract and the extent to which it presents new and significant work. The Review Committee strongly recommends that abstracts have an organized, coherent, well-thought-out, and complete presentation. Please note that no paper submitted for consideration will be eligible if it has already been, or is in consideration for, publication elsewhere at any time prior to the meeting. The winner(s) agree to publish their outstanding case report(s) in SKINmed: Dermatology for the Clinician, an official publication of the International Academy of Cosmetic Dermatology. By submitting your paper for consideration, you give SKINmed: Dermatology for the Clinician first-rights of refusal for publication through December 31, 2018.

The applicant must be in training at the time of the Congress presentation.

All applicants will receive e-mail notice of the Resident Case Report Review Committee’s decision by May 1, 2018.

Vesna Petronic-Rosic, MD, MSc
Chair, Resident Contest Committee
Professor and Chief
The University of Chicago Pritzker School of Medicine
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Basal cell carcinoma (BCC), a subtype of nonmelanoma skin cancer, is the most frequently occurring form of skin cancer in the United States and Canada. Although it seldom results in death or metastatic disease, BCC can cause significant morbidity due to its destructive local spread. The global prevalence of BCC is increasing. BCC initially affects the epidermis and later invades deeper tissues. Standard first-line treatment of BCC usually involves local sites, but still, there are cases of BCC that cannot be appropriately treated with such modalities, including recurrences after conventional therapy; therefore other forms of treatment must be considered. Sonidegib (Odomzo®, formerly LDE225; Novartis, Basel, Switzerland) is an oral, bioavailable small molecule that acts on the smoothened (Smo) receptor antagonist. It was approved in July 2015 by the US Food and Drug Administration for the treatment of locally advanced recurrent BCC (laBCC). Clinical trials have shown sonidegib to be effective and safe for the treatment of laBCC. Vismodegib (Erivedge™; Roche, Basel, Switzerland), a treatment for metastatic BCC (mBCC) and laBCC, was approved by the Food and Drug Administration in 2012.

MECHANISM OF ACTION

The sonic hedgehog (Hh) signaling pathway transmits information to embryonic cells that is required for proper cell differentiation, proliferation, and tissue patterning. Smo is an Hh signal transducer structurally similar to G protein-coupled receptors; it relies on heterotrimeric G proteins to effectively transduce the Hh signal. Sonidegib works by blocking this transduction.

PHASE I CLINICAL DATA

The first phase I trial investigated the maximum tolerated dose and dose-limiting toxicities of oral sonidegib in 103 patients with various forms of advanced solid tumors. Dosages ranged from 100 to 3000 mg daily and 250 to 750 mg twice daily in 28-day cycles. If patients tolerated the assigned dose for at least two cycles, dose escalation was allowed. Before the first cycle, there was a single-dose 7-day pharmacokinetic run-in period to describe the pharmacokinetic profile. The maximum tolerated doses reported were 800 mg daily and 250 mg twice daily.

The most frequent dose-limiting toxicity was reversible grade 3 or 4 elevated serum creatine kinase, experienced by 18% of patients at doses ≥800 mg daily or ≥250 mg twice daily. Six of the 16 (37.5%) patients with laBCC or mBCC achieved a partial (decrease in tumor size) or complete (no tumor present) tumor response after treatment, with a complete response occurring in one patient (6.25%) with laBCC treated with a twice-daily dose of 400 mg for 4 months. A strong association between BCC tumor response and Hh pathway activation, as determined by a five-gene Hh signature reverse transcriptase polymerase chain reaction assay, was found. Sufficient tumor samples were available for nine BCC patients, with all (100%) showing five-gene Hh path activity after treatment with sonidegib (Table).

In an additional phase I trial, the maximum tolerated dose of sonidegib was investigated in patients with advanced solid tumors. Twenty-one Japanese patients (group 1) were treated with either 400 or 600 mg once daily for a median of 88 and 86 days, respectively. Twenty-four patients from Hong Kong and Taiwan (group 2) were treated with 400 mg (n=12, 76

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### NEW THERAPY UPDATE

**Table. Study Characteristics, Dosing Regime, and Efficacy Rates of Odomzo in Phase I–II Clinical Trials**

<table>
<thead>
<tr>
<th>Author</th>
<th>Regime</th>
<th>Efficacy rates</th>
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<tbody>
<tr>
<td><strong>Phase I trials (n=148)</strong></td>
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<tr>
<td>Rodon et al, 2014&lt;sup&gt;9&lt;/sup&gt; (n=103)</td>
<td>Escalating doses once (100, 200, 400, 800, 1000, 1500, or 3000 mg) or twice (250, 400, or 750 mg) daily, continuously, 28-day cycle measured at day 15</td>
<td>6.3% (1/16) complete response laBCC or mBCC 31.3% (5/16) partial response laBCC or mBCC</td>
</tr>
<tr>
<td>Minami et al, 2016&lt;sup&gt;8&lt;/sup&gt; (n=45)</td>
<td>Group 1: (n=12) 400 mg, (n=9) 600 mg; group 2: (n=12) 400 mg, (n=8) 600 mg, (n=4) 800 mg, once daily; continuously 28-day cycle for a median of Group 1: 88, 86; Group 2: 76, 89 or Group 3: 74 days</td>
<td>Group 1 (n=21): 23.8% (5/21) stable disease 71.4% (15/21) progressive disease 4.8% (1/21) lost to FU Group 2 (n=24): 41.7% (10/24) stable disease 37.5% (9/24) progressive disease 20.8% (5/24) lost to FU</td>
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<tr>
<td><strong>Phase II trials (n=775)</strong></td>
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<tr>
<td>Migden et al, 2015&lt;sup&gt;11&lt;/sup&gt; BOLT (n=230) 144 M, 86 F Median age (years): group 1, 67 (25–92); group 2, 65 (24–93)</td>
<td>Group 1: 200 mg Group 2: 800 mg Once daily on a continuous dosing schedule for up to 30 days</td>
<td>Group 1 (per central review) (n=79): 3.0% (2/66) laBCC complete response 43.9% (29/66) laBCC, 15.4% (2/13) mBCC partial response 43.9% (29/66) laBCC, 76.9% (10/13) mBCC stable disease 1.5% (1/66) laBCC progressive disease 7.6% (5/66) laBCC, 7.7% (1/13) mBCC unknown Group 2 (per central review) (n=151): 35.2% (45/128) laBCC, 17.4% (4/23) mBCC partial response 43.0% (55/128) laBCC, 65.2% (15/23) mBCC stable disease 4.3% (1/23) mBCC progressive disease 21.9% (28/128) laBCC, 13.0% (3/23) mBCC unknown</td>
</tr>
<tr>
<td>Dummer et al, 2016&lt;sup&gt;10&lt;/sup&gt; BOLT FU (n=230) Same as above</td>
<td>Same as above, 12 month FU data</td>
<td>Group 1 (per central review) (n=79): 4.5% (3/66) laBCC complete response 53.0% (35/66) laBCC, 7.7% (1/13) mBCC partial response 33.3% (22/66) laBCC, 84.6% (11/13) mBCC stable disease 1.5% (1/66) laBCC progressive disease 7.6% (5/66) laBCC, 7.7% (1/13) mBCC unknown Group 2 (per central review) (n=151): 1.6% (2/128) laBCC complete response 42.2% (54/128) laBCC, 17.4% (4/23) mBCC partial response 37.5% (48/128) laBCC, 73.9% (17/23) mBCC stable disease 0.8% (1/128) laBCC, 4.3% (1/23) mBCC progressive disease 18.0% (23/128) laBCC, 4.3% (1/23) mBCC unknown</td>
</tr>
<tr>
<td>Quinlan et al., 2016&lt;sup&gt;12&lt;/sup&gt; (n=545) 399 M, 146 F Median age (years): group 1, 62.0; group 2, 34.5</td>
<td>Pooled analysis of four patient studies; measured at 17 weeks</td>
<td>Treatment group: 341 patients (n=211, 102, 21, and 7 from the phase II pivotal study A2201, study X2101, study X1101, and study B2209, respectively) Control group: 204 healthy volunteers (n=146, 36, 16, and 6 from study A2114, study A1102, study A2108, and study A2110, respectively) QTcF steady-state concentrations (200 and 800 mg): all below 5 ms Highest mean ΔQTcF: (200 mg) -3.9 ms at week 17 pre dose and (800) 2.7 ms at 2 hours post dose</td>
</tr>
<tr>
<td><strong>Phase I + II trials (n=667)</strong></td>
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<tr>
<td>Zhou et al, 2016&lt;sup&gt;13&lt;/sup&gt; BOLT and three other trials (n=667) 408 M, 260 F Mean age (years): group 1, 65; group 2, 66; group 3, 62</td>
<td>Group 1/2: 200 or 800 mg once daily for 12 weeks Group 3 PK-CK: 100–3000 mg once daily and 250–750 mg twice daily for 12 weeks</td>
<td>PK-FAS: group 1 (n=190), group 2 (n=141) and group 3 (n=336) Week 5: Cmin vs ORR indicated no relationship between Odomzo 200 or 800 mg dose and probability of complete response or partial response No exposure-efficacy relationship indicated from the progression-free survival and time to tumor response analyses Increased exposure associated with a greater risk of grade 3 or 4 CK elevation, with lower risk in women than men when Cmin used in the model</td>
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days) or 600 mg (n=8, 89 days) or 800 mg (n=4, 74 days). The maximum tolerated dose was established as 600 mg. The dose-limiting toxicity was caused by elevated creatine kinase in both treatment groups (Table).

PHASE II CLINICAL DATA

The first phase II clinical trial (Basal Cell Carcinoma Outcomes with LDE225 Treatment [BOLT]) was published in 2015 and collected data from 58 centers across 12 countries. Patients who had either laBCC not amenable to curative surgery or radiation, or mBCC were included for review and received 200 or 800 mg of sonidegib once daily (n=230). Primary results showed that sonidegib 200 mg once daily (group 1) produced a complete response in 2 of 66 (3.0%) and a partial response in 29 of 66 (43.9%) participants with laBCC; 800 mg once daily (group 2) produced a complete response in 2 of 13 (15.4%) patients with mBCC, a partial response in 45 of 128 (35.2%) for laBCC, and a partial response in 4 of 23 (17.4%) for mBCC. The 12-month follow-up data for group 1 showed a complete response in 3 of 66 (4.5%) patients, a partial response in 35 of 66 (53.0%) for laBCC, and a partial response in 1 of 13 (7.7%) for mBCC. Data for group 2 showed a complete response in 2 of 128 (1.6%) patients, a partial response in 54 of 128 (42.2%) for laBCC, and a partial response in 4 of 23 (17.4%) for mBCC (Table).

In an additional phase II trial, pharmacokinetic-pharmacodynamic analysis was performed to test whether prolongation of the QT/QTc interval occurred during extended use of sonidegib. Results showed there was no QT prolongation associated with sonidegib. No clear exposure and efficacy relationship was found with sonidegib in an additional analysis. There was, however, a positive relationship between sonidegib exposure and grade 3 creatine phosphokinase elevation. Comparing estimated steady-state concentration minimums, 800 mg once daily was found to deliver an exposure level 2.3-fold higher than a dose of 200 mg once daily (Table).

![Figure: Adverse events of Odomzo reported in phase I–II clinical trials.](image_url)
CONCLUSIONS

Sonidegib has, on the strength of the Phase II data, been approved by the Food and Drug Administration for the treatment of laBCC recurring after surgery or radiation therapy, and for individuals who are not candidates for surgery or radiation therapy. At the time of writing, no phase III trials involving sonidegib have been published. There is a ‘black box’ warning for sonidegib for embryonic/fetal toxicities.7 Adverse events associated with sonidegib were mostly grade 1 (mild) and grade 2 (moderate) in severity (eg, nausea and fatigue) (Figure 1). A malfunction of the Hh signaling pathway involving one or both P53 and MC1R gene mutations increases the risk of laBCC development; thus medical management with Smo receptor antagonists, such as oral sonidegib and vismodegib, seems to be an effective strategy against the progression of laBCC.14

The approval of sonidegib adds to the armamentarium of treatments for recurrent laBCC that could improve patient quality of life. There is a lack of head-to-head, long-term studies comparing sonidegib to its chief competitor vismodegib and to other oral treatments approved for the treatment of laBCC, and such studies would be valuable. Sonidegib may be an option for patients with laBCC and for physicians searching for a nonsurgical alternative or additional therapy for this disease.

REFERENCES

Malignant spindle cell tumors are typically diagnosed by biopsy, often confirmed with immunohistochemical staining, and treated by excision with wide margins; however, the more benign variants of spindle cell tumors do not require such aggressive treatment. These include leiomyomas, angioleiomyomas, neurofibromas, schwannomas, neurothekeomas, hemangiomas, glomus tumors, myofibromas, and spindle cell lipomas. Immunostaining is recommended to confirm the benign nature of these tumors, not only due to the difference in treatment of benign and malignant spindle cell tumors, but also because other malignant tumors may also have spindle cell components or even be disguised as spindle cell tumors.

AN ILLUSTRATIVE PATIENT

Over a 3-year period, a 57-year-old man developed a soft, nodular, asymptomatic, flesh-colored lesion on the skin over the lateral aspect of the left clavicle. The lesion was focally ulcerated, but no telangiectasia was noted. The lesion was initially diagnosed, based on histopathologic presentation on hematoxylin and eosin staining, as a spindle cell tumor of uncertain differentiation and uncertain behavior (Figures 1 and 2). Subsequently, BerEP4 staining identified the lesion as a basal cell carcinoma (BCC) (Figures 3 and 4). The patient was treated appropriately for a malignant tumor by excision with wide margins.

DISCUSSION

BCCs have been previously reported to contain histologic areas with a proliferation of spindle cells. A rare entity, known as sarcomatoid BCC, may contain spindle cell and basal cell components. The spindle cell component can be homologous or heterologous. Previous reports have suggested that investigating these tumors under low magnification may lead to missing the BCC; higher power magnification and confirmatory staining can highlight typical features of BCC. As this particular variant has aggressive behavior and metastatic potential, prompt recognition of the malignant component is important. In addition, there may be spindle cell tumor and BCC in the same lesion. One report described a “hybrid” tumor with characteristics of BCC and myoepithelioma. The spindle cells stained strongly with p63. Neurofibroma and BCC have also been reported to occur in the same lesion, presenting clinically as nonulcerating nodules.

Spindle cell squamous cell carcinoma is uncommon but not rare. This variant is a relatively aggressive tumor often found in patients with a history of radiation burn, scars, or organ transplantation; it has the potential for local recurrence and distant metastases. Histologically, this tumor has a spindle cell component that infiltrates the dermis as individual cells or nests of cells. The spindle cells are interspersed with varying degrees of squamous cell carcinoma features. This tumor usually stains positive for 34βE12, p63 and low-molecular-weight keratin (AE1/AE3).

Multiple theories for the presence of two histologic subtypes in the same lesion have been proposed. The collision theory states that two independent tumors may coincidentally collide in the same location, explaining the presence of both histologic subtypes. This theory may account for some cases, but it is not the most widely accepted explanation. The composition theory states...
Basal or Squamous Cell Carcinoma Mimics a Spindle Cell Tumor

**Figure 1.** Case 1: basal cell carcinoma resembling a spindle cell tumor (hematoxylin and eosin stain, magnification ×270).

**Figure 2.** Case 1: basal cell carcinoma resembling a spindle cell tumor; ulcerated area. (hematoxylin and eosin stain, magnification ×540).

**Figure 3.** Case 1: BerEP4 stain identifying the lesion as a basal cell carcinoma (BerEP4 stain, magnification ×270).

**Figure 4.** Case 1: BerEP4 stain confirming basal cell carcinoma, showing the cellular detail (BerEP4 stain, magnification ×540).
that the spindle or stromal cell component of the tumor is actually a reaction to the malignant epithelial component. The combination theory suggests that both components arise from a common progenitor cell that undergoes divergent differentiation. Finally, the conversion theory suggests that the spindle cell component arises from a metaplastic transformation of the epithelial component. This last theory is most widely accepted due to genetic evidence of monoclonality in lesions, including both histologic subtypes, showing identical genetic mutations of p53 and k-ras, loss of heterozygosity, and similar genetic alterations.

Although it is appropriate to be conservative in the treatment of benign spindle cell tumors, dermatologists should be aware of the occasional malignant tumor that may be hidden amidst the bland ocean of spindle cells. We recommend confirmatory staining of spindle cell tumors for two reasons:

• first, as an aid in definitively differentiating a benign spindle cell tumor from its malignant counterpart, especially as the clinical differentiation is often unclear;

• second, because, while rare, there a potential for multiple histologic subtypes to exist in the same lesion.

CONCLUSIONS
When the clinical-pathologic correlation is not clear, we recommend investigation for another diagnosis, such as BCC or squamous cell carcinoma.

REFERENCES
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The Prescience of Bernie Ackerman: The Road from “Pseudoleukemia Cutis” to “Intralymphatic Proliferation of T-cell Lymphoid Blasts (IPTCLBs)”
Warren R. Heymann, MD

“I have been accused, perhaps justifiably, of having written more than I have read. I would not presume to speak for the readers of my writings, but I can speak for myself: How enormously instructive and pleasurable it has been! I hope that readers feel the same.” —A. Bernard Ackerman

As with anyone’s passing, life goes on and memories fade. Periodically, there is a jolt—a reminder of a person’s brilliance and legacy left behind. I never trained directly with Bernie but had the good fortune to attend his lectures and become acquainted with him personally. There is no dermatopathologist alive who has not been influenced by his teachings, whether they agree with him or not. Based on the quote above, I can assure him posthumously that his writings are still incisive and instructive.

In 1977, prior to the advent of immunohistochemistry, Ackerman and Tanski described “pseudoleukemia cutis.” The following is the abstract from the paper:

Histologic sections from a solitary cystic cutaneous lesion that showed atypical mononuclear cells in the dermis and within blood vessels were diagnosed by several general pathologists and dermatopathologists as leukemia cutis. The patient, who had no other cutaneous lesions, was consequently submitted to an extensive investigation for leukemia, which proved negative. Additional and deeper sections from the original block revealed that the cellular infiltrate so suspicious of leukemia cutis was secondary to rupture of a lesion of molluscum contagiosum. The correct histopathologic diagnosis, therefore, was pseudoleukemia cutis. The lessons of the case are that 1) further study of the specimen, solitary as it was and asymptomatic as the patient was, would have obviated worry and the expense and inconvenience of an extensive systemic investigation, and that 2) the diagnosis of leukemia cutis should never be made solely on the basis of histologic sections of skin, but rather after examination of blood and bone marrow.

In the photomicrographs in their paper, one can clearly see atypical mononuclear cells within vessels. It is certainly understandable why seasoned pathologists thought this patient had leukemia.

INTRALYMPHATIC PROLIFERATION OF T-CELL LYMPHOID BLASTS (ILPTCLBS)

Fast forward four decades to the increasingly reported concept of ILPTCLBs—intralymphatic proliferation of T-cell lymphoid blasts. The most recent report is that of a 77-year-old man who was treated with antibiotics for a hip prosthesis infection. Eight weeks later, he developed an eruption characteristic of DRESS (drug reaction with eosinophilia and systemic symptoms) syndrome, accompanied by fever and eosinophilia. Cutaneous biopsy revealed an atypical T-cell proliferation within dermal lymphatic vessels. The lymphocytes were mid-sized, with mitoses and apoptotic figures. They were CD3+, CD4+, CD5+, and some were CD30+. There was no T-cell receptor clonal rearrangement. Complete regression of the cutaneous eruption and eosinophilia was observed after ceasing treatment with antibiotics. The diagnosis was that of a benign atypical intralymphatic

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T-cell proliferation associated with DRESS syndrome. The authors assert that the occurrence of atypical dermal CD30+ T cells in cutaneous biopsy during benign reactive conditions such as arthropod bites or scabies is well known. The intralymphatic localization of atypical reactive lymphocytes, however, is much less common and represents a diagnostic pitfall because it can suggest an aggressive intravascular lymphoma.3

ILPTCBs have also been noted in other inflammatory disorders such as hidradenitis suppurativa4 and lichen sclerosus.5 The histopathologic criteria for the diagnosis of ILPTCBs are: (1) the presence of an associated inflammatory disorder; (2) clusters of large atypical lymphoid cells confined to D2-40+ lymphatic vessels, although scattered atypical extravascular cells are common; (3) a T-cell phenotype without aberrant features (no loss of pan-T-cell markers, no aberrant double positivity/negativity of CD4 and CD8); (4) EBER-1 negativity for the Epstein-Barr virus; and (5) a polyclonal infiltrate as detected by polymerase chain reaction.4

THE DIFFERENTIAL DIAGNOSIS OF ILPTCBs

Intralymphatic histiocytosis enters the histologic differential diagnosis, and is classically associated with rheumatoid arthritis.3 Immunohistochemistry clearly differentiates intralymphatic histiocytosis from ILPTCBs; in intralymphatic histiocytosis, the infiltrate is positive for CD68 and CD163, in contrast to the T-cell profile observed in ILPTCB.4 Of course intravascular lymphoma or leukemia must always be ruled out. Intravascular lymphoma or leukemia is an aggressive disease, with most cases representing a diffuse large B-cell lymphoma, although a rare T-cell variant has been reported, including cases with CD30 expression.5 Recently, an indolent form of intralymphatic cutaneous anaplastic large cell lymphoma/lymphomatoid papulosis, with CD30 positivity, has been reported.6

CONCLUSIONS

When intralymphatic atypical lymphocytes are appreciated, it is essential to look at the clinical context, perform immunohistologic stains, and check for clonality, in order to differentiate a benign process from a true lymphoma. As Bernie taught us, clarity in the writing and linguistics of dermatopathology is essential to translate what is seen microscopically to useful clinical information. I am not sure that a report of ILPTCB would mean anything to the referring dermatologist without a detailed explanation. Perhaps we would be best utilizing Bernie’s original term, “pseudoleukemia cutis.”

REFERENCES

A 23-year-old woman presented with asymptomatic, flesh-colored papules of 8 years’ duration on the chin. She gave a history of recurrent erythematous papules and pustules that used to subside in 2–3 weeks. On examination, multiple, grouped, flesh-colored papules along with a few erythematous papules were noted on the chin (Figure 1). The rest of the mucocutaneous examination was unremarkable except for a few erythematous papules around the nose.

Histopathology of a specimen taken from a flesh-colored papule showed numerous hypertrophied sebaceous glands with extensive surrounding fibrosis. A superficial and mid-perivascular and periappendageal lymphoplasmocytic infiltrate were noted (Figure 2). On clinicopathological correlation, a diagnosis of gnathophyma was made.

Rhinophyma is a well-known complication of rosacea, but phymas of other regions (chin, helix, forehead, eyelids) are uncommon and can cause confusion in diagnosis. The phymatous changes affecting areas other than the nose are usually infrequent and do not occur as an isolated entity.1 Our patient had almost exclusive involvement of the chin—a rare presentation.

Figure 1. Grouped, flesh-colored papules on the chin.

Figure 2. (A) Numerous hypertrophied sebaceous glands with a perivascular and periappendageal lymphoplasmocytic infiltrate (hematoxylin and eosin, original magnification ×40). (B) Hypertrophied sebaceous glands with extensive surrounding fibrosis (hematoxylin and eosin, original magnification ×400).

REFERENCE

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Robert S. Berger, MD

JOINT SESSION WITH THE INTERNATIONAL SOCIETY OF DERMATOLOGY: BUENOS AIRES, ARGENTINA, APRIL 18

Richard Spielvogel, MD (Philadelphia, PA) described horizontal sectioning of hair biopsies as the best way to obtain a correct diagnosis of hair disorders. The average hair count ranges up to 24 per section for this method, as opposed to four for vertical sections. Vertical sections may, however, be required to visualize the dermoeidermal junction.

Anthony V. Benedetto, DO (Philadelphia, PA) discussed avoiding postoperative surgical site infections by promoting wound healing with a topical silicone gel that is semicocclusive, gas-permeable, and bacteriostatic. When applied to an open or closed wound, the agent forms a flexible, protective film barrier that contours to the wound surface, reducing transepidermal water loss and inflammation.

Dee Murell, MD (Sydney, Australia) noted a 50-fold increased incidence of squamous cell carcinoma in recessive dystrophic epidermolysis bullosa (EB), which appears well differentiated but has a very poor prognosis with early metastasis. Squamous cell carcinoma can occur at a very young age (the youngest case described being 7 years old) and in all skin types, including non-sun-exposed and recurrent trauma sites. Recessive dystrophic EB patients with squamous cell carcinoma have a much higher level of urinary basic fibroblast growth factor than individuals with other types of EB, or recessive dystrophic EB patients without squamous cell carcinoma.

BARILONCHE, ARGENTINA, APRIL 24

Allan Arbuckle, MD (Denver, CO), in his update on EB, reported that a skin biopsy is now sufficient instead of a kidney biopsy in diagnosing Alport syndrome. The most debilitating symptoms of EB simplex are hyperkeratosis of the hands and feet. Junctional EB with granulation tissue in the middle region of the face carries a poor prognosis, whereas patients with nonlethal forms develop nail abnormalities. Kindler syndrome shows skin fragility and photosensitivity with actinic keratoses detected in children as young as 5 years old.

Anthony V. Benedetto, DO, discussed use of botulinum toxin A in the lower region of the face, where the angle of the chin can be blunted. When injecting the platysmal band, the 2-unit injections should be administered very superficially.

A. Howland Hartley, MD (Leonardtown, MD) presented newly reported galactose-α1,3 galactose sensitivity, the first reported allergy to a nonprotein. It had been described in patients taking cetuximab, and is now seen in the general population. There is a delayed reaction to the nonprotein allergen, which is found in mammalian meats and transmitted by a Lone Star Tick bite.

Isabel Maria del Pilar Casas, MD (Patagonia, Argentina) presented a wonderful “how to” approach for educating school age children in Argentina about sun
damage and protection. Because a child's attention span is about 7 minutes, brevity and interaction are critical, and the use of graphic photos should be avoided. See the excellent YouTube videos “The Riddles of the Sun” (in Spanish with English subtitles; https://www.youtube.com/watch?v=mzaHSa8ghKw) and “Acertijos del sol” (in Spanish; https://www.youtube.com/watch?v=YC85bjbMtiw).

CALAFATE, ARGENTINA, APRIL 26–27 (FIGURE)

John R. Hamill, MD (Tampa, FL) discussed the new use for tazarotene cream. Under occlusion, this retinoid can be used for treating squamous cell carcinomas that have developed on the legs as a result of radiation therapy for lichen planus.

H. Alan Arbuckle, MD (Denver, CO) emphasized that the most important component of wound care is treating the underlying cause. With moisture balance and protein balance being critical, necrotic tissue should be removed frequently, as the colonizing bacteria can create problems. Vaseline or occlusive dressings are preferred over wet-to-dry dressings.

Dedee Murrell, MD told the group about a newly discovered defect to the 19th gene (currently named KLHL24), which causes EB and is involved in degradation of keratin 14. The first gene therapy for a skin disease has been performed in junctional EB; this involved stem cells, using a retroviral vector grown into artificial skin equivalents.

Robert Bryg, MD (Olympia, WA) reminded the group that electronic medical records are not a panacea. Errors, uncommon abbreviations, cloning of notes, and just plain carelessness can lead to payment denial.

Paul Garson, MD (Hampton, NY) noted that the addictive substances of opium, alcohol, and cocaine have been present in society for hundreds of years. Addiction involves the pleasure center of the brain, where excitement about the expectant pleasure resulting from use leads to increased dopamine release, flooding the brain and producing the resulting pleasure. Tolerance ensues, requiring more substance use and then addiction.

GALAPAGOS ISLANDS, MAY 2

Robert S. Berger, MD (White Plains, MD) reported that the cost of drugs has worsened. Within the previous year, the price of generic medication had sky-rocketed (clobetasol being the most recent example). With respect to branded medications, Targretin (bexarotene) has been one of the more egregious examples of overpricing, increasing to almost $31,000 per 60 g tube (with no generic available) and $51,000 per 200 capsules (oral). A new generic bexarotene tablet costs $24,000 to $31,000 per 200 (as per https://www.goodrx.com price listings). Independent specialty pharmacies are making inroads in cost reductions, and some US states are considering legislation to cap drug costs.

ANNUAL MEETING APRIL 10–20, 2018

The 59th Annual Meeting will take place in New Zealand, with first sessions in Auckland on North Island, followed by Christchurch on South Island, April 10–20, 2018.
It seemed like a good idea at the time. The *Phylloxera* pest was devastating Hungary’s famous vineyards, and bootleg lower quality wines were ruining what was left of the region’s reputation. To detect adulteration, in 1893 the Hungarian Parliament mandated that all wines contain the additive phenolphthalein. Shortly thereafter, a nationwide epidemic of diarrhea broke out, demonstrating the hitherto unknown fact that phenolphthalein was not only an excellent acid-base indicator, but also a powerful purgative.

One of the first to profit from this serendipitous discovery was a Budapest pharmacist, Dr. D. Bayer. Although phenolphthalein itself is tasteless, he decided to enhance its appeal by adding sugar and vanilla flavoring. Dr. Bayer’s laxative, marketed under the all-too-descriptive name Purgen, was an immediate success. Shortly after its introduction in 1900, a host of copycat vanilla-flavored laxatives flooded the market.

Considering the popularity of Purgen® and its numerous imitators, it should come as no surprise that a fellow Hungarian pharmacist was responsible for the next innovation in laxatives. Max Kiss had immigrated to Brooklyn, NY, in 1905, bringing with him the idea of a chocolate-flavored phenolphthalein laxative. The original name for his brainchild, BoBo, was quickly rejected, when he learned this word was also vulgar street slang. In a Hungarian language newspaper, Kiss read about the political deadlock in his homeland. The term for this constitutional stalemate was “ex-lex.” Eureka! Kiss christened his invention Ex-Lax®, shorthand for Excellent Laxative.

On August 29, 1997, the Food and Drug Administration proposed a ban on phenolphthalein. Recognizing that further resistance was futile, the next day the makers of Ex-Lax® voluntarily removed their product from store shelves. Ex-Lax® was reformulated, with senna taking the place of phenolphthalein. Given it was phenolphthalein that triggered the allergic reaction, fixed drug eruptions due to Ex-Lax® passed into history.*

REFERENCES


*Speaking of historical footnotes, Dr. Fox’s patient was the first reported case of laxative-induced fixed drug eruption.

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W W W. T N L A S E R S O C I E T Y . C O M
CASE STUDY
Vesna Petronic-Rosic, MD, MSc, Section Editor

Coccidioidomycosis
Matthew Laffer, Lindsay Ackerman, MD

A 23-year-old man presented with new-onset pruritic and painful urticarial lesions and targetoid erythematous plaques on both palms, the trunk, and the upper and lower extremities (Figure 1). Additionally, small pustules were discovered on the neck (Figure 2), and there with edematous erythematous vermillion lips with splaying onto the cutaneous lips without ulceration. The patient stated he had had a fever before the eruption, fatigue, chills, myalgias, and sore throat. A chest x-ray was obtained and showed bilateral infiltrates. Two 4-mm punch biopsies were performed on the left forearm and left side of the neck; a resulting section from the left forearm is shown in Figure 3. (SKINmed. 2018;16:51–53)

Histologic examination of the biopsy specimens revealed spongiotic dermatitis, one of the few patterns often seen in cutaneous eruptions in patients with pulmonary or systemic coccidioidomycosis. Periodic acid–Schiff staining was negative for fungal organisms.

A complete blood count with differential revealed a neutrophil leukocytosis, and results of a comprehensive metabolic panel were within normal limits. Samples were negative for Mycoplasma pneumoniae IgM antibody, and positive for Coccidioides antibody.

The only significant factor in the patient’s past medical history was that previously he had suffered from herpes zoster. He stated that he was 12.5% Filipino, had been born and raised in Michigan, but had recently moved to Arizona.

The patient was treated with a 14-day course of 400 mg fluconazole orally and an oral steroid taper, and was also given triamcinolone acetonide 0.1% cream to apply twice a day, as needed. Within 1 month after completion of his medications, the pulmonary and cutaneous clinical manifestations had completely cleared.

Coccidioidomycosis (cocc) is a fungal disease that is prevalent in the southwest United States, northern Mexico, and parts of Central and South America. In the United States, the incidence is estimated to be 100,000 cases per year. This disease is caused by the dimorphic, highly virulent pathogen Coccidioides immitis; infectious arthroconidia live in the soil and are inhaled.

Approximately 60% of patients have an asymptomatic infection, while most others have a self-limiting pulmonary infection. Extrapulmonary complications vary, and the patient’s ancestry and immune status may play a large role. In individuals of Caucasian ancestry, extrapulmonary complications are estimated to occur in as few as 0.5% of infections, whereas in individuals of African or Filipino ancestry, they may be significantly more common. Extrapulmonary complications may occur in as many as 50% of immunosuppressed patients, including those with AIDS or lymphoma, and those using high-dose corticosteroids. Extrapulmonary dissemination most frequently involves the skin, but commonly involves the meninges, bones, joints, and soft tissues. Patients that present with clinical manifestations of an acute respiratory illness often display chest pain, cough and fever.

Cutaneous manifestations of coccidioidomycosis are variable, and can be classified as organism-specific or reactive. Organism-specific infections result from hematogenous spread to the skin in disseminated disease, or very rarely from primary cutaneous inoculation. These plaques often present as “lesions,” morpho-

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logically represented as papules, nodules, abscesses, sinus tracts, or large verrucous plaques. The histopathology of a primary lesion demonstrates dense infiltrates of neutrophils, lymphocytes, and plasma cells, at times with necrosis, microabscess formation, or granulomas, with evidence of cocci spherules.

Reactive cutaneous manifestations of cocci are clinically variable and can present as toxic erythema, erythema multiforme, morbilliform exanthems, erythema nodosum, and Sweet-like reactions. Commonly, the histology of cutaneous reactions to cocci infections includes perivascular lymphocytic infiltrates with neutrophils and eosinophils in the dermis, and epidermal spongiosis that can be mild or very marked, causing basilar keratinocyte vacuolization. Because the histology is variable, serologic tests and fungal cultures are helpful to confirm the diagnosis.

CONCLUSIONS

Treatment of a coccidioidomycosis is dependent upon the extent of pulmonary compromise, the presence of dissemination, and underlying patient risk factors. Treatment for uncomplicated pulmonary cocci is controversial, whereas azole antifungals are used for severe pulmonary infections, disseminated disease, or infections occurring in immunocompromised individuals.
REFERENCES


HISTORICAL DIAGNOSIS AND TREATMENT

Diagnosis and treatments have advanced over the past century. This feature depicts conditions from a collection of stereoscopic cards published in 1910 by The Stereoscopic Skin Clinic by, Dr S. I. Rainforth.

(Continued on page 75)
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CASE STUDY

Tuberous Sclerosis Complex: A Classic Presentation

Virendra N. Sehgal, MD, FNASc, FAMS, FRAS (Lond);¹ Navjeeven Singh, MD;² Sonal Sharma, MD;² Jolly Rohatgi, MS;³ Rakesh Oberai, MD;⁴ Kingshuk Chatterjee, DNB, MNAMs⁵

An 11-year-old girl presented with an insidiously evolving, reddish-brown, small, hard, elevated lesion, occupying the midsection of her face, which had been present since early childhood. There were also a few small white spots over the trunk. There was no history of seizures or visual deficit, and no burning on exposure to sunlight. There were no known congenital defect noted at birth, and her parents were nonconsanguineous. There was no significant family history. There were numerous 2- to 4-mm reddish-brown papules located symmetrically on the nose, nasolabial folds, and cheeks (Figure 1A). In addition, there was an uneven 3-cm plaque in the lumbosacral region that resembled orange peel—a shagreen patch (Figure 1B). There were also two well-defined, 5- to 10-mm, hypomelanotic, ivory-white macule(s) with irregular margins (Figure 1C). The buccal mucosa and nails were unremarkable, and indirect ophthalmoscopic and slit-lamp examination of the eye was normal. Laboratory studies were unremarkable. Ultrasonography of the abdomen was normal, as were abdominal and chest x-rays. (SKINmed. 2018;16:55–58)

Hematoxylin and eosin–stained sections prepared from the hard plaque on the face revealed a normal-looking epidermis, small pilosebaceous units, and evidence of increased dermal collagen (Figure 2A). At higher magnification, “onion-peel”-like concentric bands of collagen were seen surrounding abortive hair follicles (Figure 2B). Concentric bands of fibrosis were also seen around some of the eccrine ducts (Figure 2C), appearing to constrict their lumina. The small sebaceous glands in the dermis were surrounded by dense collagen bands.

Sections from the plaque on the back showed a dome-shaped elevation of the epidermis caused by dense fibrous tissue, and a paucity of dermal appendages and vasculature. Concentric bands of collagen, surrounding small, congested blood vessels, were also seen in the dermis.

MAGNETIC RESONANCE IMAGING (MRI)

MRI of the brain was performed on a 1.5 T MRI scanner (Signa HDxt; GE Medical Systems, Milwaukee, WI) using turbo spin-echo T1-weighted, T2-weighted, and fluid-attenuated inversion recovery (FLAIR) sequences in the axial plane. This showed subependymal nodules along the walls of the lateral ventricles, which appeared hyperintense on T1- and T2-weighted MRI (Figure 3). There were also small focal lesions in the left frontal lobe cortex (Figure 4) and left cerebellum (Figure 5), which appeared hyperintense on T2-weighted and FLAIR images, and isointense to hypointense on T1-weighted images (Figure 5). In addition, bilateral bandlike hyperintense lesions were seen in the deep white matter on FLAIR images. An extra-axial cystic lesion was also seen in the left temporal region, appearing hypointense on FLAIR and hyperintense on T2-weighted images. The orbits were unremarkable.
Ultrasonography of the abdomen, including the kidneys and liver, was normal.

**DISCUSSION**

Tuberous sclerosis is a genetic disorder in which cellular differentiation and proliferation results in hamartoma formation in the skin, brain, eye, kidney, and heart.\(^2\) Friedrich Daniel von Recklinghausen (1833–1910)\(^3\) provided the initial description in 1862, while Désiré-Magloire Bourneville\(^4\) (1840–1909) coined the term *sclérose tubéreuse*, from which the current name...
Figure 3. (A) Axial T2-weighted image demonstrating hyperintense subependymal nodules (arrows) along the lateral ventricles. (B) On axial T1-weighted imaging, the nodules appeared hyperintense with central hypointensity (arrows).

Figure 4. Axial fluid-attenuated inversion recovery images demonstrating hyperintense linear radial bands in the deep white matter bilaterally (white arrows) and a focal hyperintense lesion in the left parietal cortex (black arrow).

Figure 5. Axial T2-weighted image showing a focal hyperintense lesion in the left cerebellum (thin arrow) and an extra-axial hyperintense lesion in the left temporal region (thick arrow). (B) On axial fluid-attenuated inversion recovery images, the left cerebellar lesion appeared hyperintense (thin arrow), and the extra-axial left temporal lesion appeared hypointense (thick arrow).

Table. Tuberous Sclerosis Complex: Criteria for Diagnosis

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<td>• Multiple renal cysts</td>
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<td>• Multiple retinal nodular hamartomas</td>
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<td>Tumors in other organs</td>
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<td>• Cardiac rhabdomyomas</td>
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<td>• Lymphangioleiomyomatosis</td>
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<td>• Renal angiomyolipomas</td>
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To avoid “double-counting” because central nervous system lesions can also cause seizures, the criteria do not include seizures/intellectual disability.

Our patient had facial angiofibromas, a shagreen patch, and hypomelanotic, confetti-like skin lesions as a few of its salient cutaneous manifestations, which was adequate to consider the diagnosis of TSC as it conformed to the major features of this condition (see above).

The microscopic pathology was significant enough to warrant recommending a skin biopsy before obtaining an MRI scan. Multiple high-signal MRI lesions are characteristic of TSC, corresponding to the hamartomas and gliotic area seen patho-
logically. MRI may be useful in predicting the eventual clinical severity in younger children with newly diagnosed TSC.

Cortical tubers, subependymal nodules, and white matter lesions are commonly encountered in TSC. Less common manifestations include cerebellar atrophy, dysgenesis of the corpus callosum, Chiari malformations, arachnoid cysts, and infarctions due to occlusive vascular disorders. MRI is the most sensitive modality for their detection.\textsuperscript{13,14}

Cortical tubers, subependymal nodules, and white matter lesions, and a left temporal arachnoid cyst were seen in the present case. The cortical tubers showed increased signal intensity on T2-weighted and decreased signal intensity on T1-weighted sequences.

On MRI, subependymal nodules are isointense/hyperintense on T2-weighted images and hyperintense on T1-weighted images, and may also demonstrate central low T2 signal with a surrounding hyperintense rim. Subependymal nodules typically measure less than 1 cm but may grow over time, giving rise to subependymal giant cell astrocytomas, which are commonly located at foramen of Monro, and can cause acute obstructive hydrocephalus. They are heterogenous on MRI, contain calcification, and show intense unhomogeneous enhancement. Superficial white matter abnormalities—radial white matter bands, which are lines of arrested neuronal migration—appear as radiating bright lines on T2-weighted and FLAIR images. White matter cystic lesions may also occur.

MRI, therefore, plays an important role in arriving at the correct diagnosis of TSC, especially in asymptomatic patients, to determine the presence and extent of organ involvement. Follow-up MRI should be carried out for evaluation of the lesions and early detection of associated complications.

**CONCLUSIONS**

Although TSC is a well-known genetic disorder, there are only sporadic reports describing it. Slowly evolving angiofibromas affecting the mid-facial region, a shagreen patch, and hypomelanotic, confetti-like lesions should arouse suspicion. The clinical diagnosis should be supplemented by histopathology. Ultimately, carrying out an MRI to define the neurologic findings, plus a slit-lamp examination, is recommended.

**REFERENCES**

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A 58-year-old man presented to our department with multiple chronic vegetant plaques on the face and scalp that had been present for 4 months. The plaques were well-defined hypertrophic areas, measuring between $3 \times 3$ cm and $3 \times 4$ cm, that were located mainly on the face symmetrically and on the forehead (Figure 1). The patient did not report itching or pain. There were also dry, crusty, well-defined plaques on the scalp. No other lesions were present on the skin or mucous membranes. (SKINmed. 2018;16:60–61)
We report here a patient with the unusual presentation of dry vegetating lesions confined to the face and scalp. In nonintertriginous locations, as in our patient, the vegetating plaques dry out, evolving into warty, fissured, painful plaques that, on the face, simulate seborrheic keratoses; hence, it is classified as dry pemphigus vegetans.

The diagnosis of pemphigus vegetans is based on clinical manifestations and confirmed by histology. Lesional histopathology typically shows papillomatous, verrucous, or pseudoepitheliomatous hyperplasia. Suprabasal acantholysis can be masked by an exuberant proliferation of involved epidermis. Intraepidermal microabscesses consisting of eosinophils and neutrophils are often seen. It is associated with antibodies against desmogleins, mainly desmoglein 3. In this rare type of pemphigus, it appears that the different pathogenicity of the autoantibodies plays a role in the peculiar clinical presentation of affected patients.

The main clinical differential diagnosis of facial pemphigus vegetans with similar clinical manifestations is seborrheic keratosis, but this was ruled out here by the skin biopsy. In addition, an incorrect diagnosis of pemphigus erythematosus could have been made due to the localization of the crusty plaques in the seborrheic areas. This diagnosis was also ruled out by histologic findings consistent with suprabasal acantholysis.

The treatment of pemphigus vegetans is the same as for pemphigus vulgaris. Systemic corticosteroids are the mainstay of therapy and should be combined with immunosuppressive therapy such as azathioprine, as in our patient. The overall prognosis is good, with a low mortality rate. Patients with Neumann type pemphigus vegetans may show a severe disease course similar to that of pemphigus vulgaris.

CONCLUSIONS

The present case was unusual, as the lesions were nonflexural in distribution, and the oral mucosa was not involved. A high index of suspicion is required to diagnose pemphigus vegetans at nonintertriginous sites.

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Routine blood studies and HIV screening were negative, as were a chest X-ray, a computed tomography scan of the brain, and an abdominal ultrasound. The lesions were surgically excised with electrodessication of the bases. Histopathologic examination of the skin lesions showed multiple sporangia and spores with morphologic features of *Rhinospo-
ridium* species along with foreign body giant cells in the der-
mis (Figure 2). When a final diagnosis of nasal-nasopharyngeal rhinosporidiosis with cutaneous dissemination had been made, the patient was started on oral dapsone (100 mg per day) for 3 months. He improved and did not show any recurrence of the nasal or cutaneous lesions.

**DISCUSSION**

Rhinosporidiosis is a chronic recurrent granulomatous dis-
ease of humans and animals, caused by *Rhinosporidium seeberi*. It is endemic in some areas of Asia, such as south India and Sri Lanka, but infections have been reported in the Americas, Europe, and Africa. In India, the disease is endemic in Chhattisgarh, Tamilnadu, Kerala, parts of Orissa, and eastern Madhya Pradesh. The causative organism was first reported by Malbran but later described by Guillermo Seeber, both from Buenos Aires, Argentina, giving it its name, *R. seeberi*.

The taxonomy of the causative agent of rhinosporidiosis has always been controversial. Seeber considered the sporangium of *R. seeberi* to be a sporozoon. Although the agent was con-
sidered to be a fungus, it was interpreted as a protozoan parasite, a cyanobacterium, and a carbohydrate waste product. It has been proposed that the organism should be considered in a new eukaryotic group of protists known as Mesomycetozoa. Others have agreed with the concept of a novel clade of aquatic protistan parasites named Ichthyosporea. The causative agent of rhinosporidiosis may well be the cyanobacterium *Microcystis aeruginosa*, isolated from clinical samples as well as from samples of water in which patients have been bathing. The life cycle of the parasite is complicated. The mature forms of the organism, known as sporangia, contain multiple sporangiospores. The trophocytes, the immature forms of *R. seeberi*, are smaller and thinner than sporangia and do not contain endo-
spores. Sporangiospores are released at maturity and thereafter develop into trophocytes. The disease is possibly transmitted to humans by direct contact with spores through dust, infected clothing, or fingers, and by swimming in stagnant waters. Al-
though a large number of people living in endemic areas bathe in common ponds, only a few develop the disease. This indicates the existence of predisposing factors in the host.

Blood group studies indicate that rhinosporidiosis is common in patients with group O and group AB blood; however, the blood group distribution is too variable to draw any conclusion. Goihman-Yahr has postulated that a lack of digestive capacity of
phagocytes from affected patients may explain why only a few individuals acquire the infection.6

Clinically, morphology of rhinosporidiosis lesions is characterized by the development of pedunculated and sessile polypoid lesions on the mucosa of the nose, eyes, and larynx, and very rarely on other parts of the body, such as the skin, viscera, and brain. Disseminated cutaneous lesions are very rare and are generally associated with mucosal lesions. Cutaneous lesions without mucosal involvement are extremely rare.7,8

Three main types of lesions appear on the skin:

1. satellite lesions around the nose in cases of nasal rhinosporidiosis;
2. generalized skin involvement with or without nasal involvement caused by hematogenous dissemination;
3. primary cutaneous lesions, which may occur by direct inoculation of the organism into the skin7

Rhinosporidiosis has a multifaceted presentation on the skin. Cutaneous lesions have been reported as pedunculated or sessile growths, verruca vulgaris–like lesions, friable nodular lesions, subcutaneous swellings, furunculoid lesions, cutaneous horns, shiny globular swellings, cutaneous ulcerations, and cystic swellings. Development of a cutaneous lesion may be an indication of early
dissemination, and a thorough search should be made to exclude systemic involvement; our patient did not reveal any systemic lesions. The various clinical differential diagnoses include verrucae vulgaris, verrucous tuberculosis, and granuloma pyogenicum.7

As the organism cannot be grown in culture, histopathology is the gold standard.9 Giemsa imprinted smears and a fine needle aspirate with 10% KOH examination are also helpful.7 A biopsy may reveal a hyperplastic epithelium with a chronic inflammatory cell infiltrate composed of plasma cells and lymphocytes along with foreign body giant cells. Characteristic sporangia in various stages of maturation are seen as globular cysts of various sizes (50 to 1000 μm in diameter), lined by a well-defined wall and containing numerous endospores of diameter 5 to 10 μm. Sporangia and endospores stain positively with various special stains such as Gomori methenamine silver, periodic acid–Schiff, mucicarmine, Grocott stain, and hematoxylin and eosin.10 The periodic acid–Schiff stain is helpful in discriminating between endospores and epithelial cells, especially from the nasopharynx, as the endospores stain markedly magenta, while the epithelial cells are periodic acid–Schiff–negative.9

Mucicarmine stain is particularly helpful in differentiating Coccioidoides immitis, as sporangia and spores of this organism do not stain positively. Coccioidiomycotic lesions can cause confusion with rhinosporidiosis during cytologic as well as histopathologic evaluation, as the former has similar mature stages represented by large, thick-walled spherical structures containing endospores. Distinction can also be made by hematoxylin and eosin stain as the intrasporangial endospores of C. immitis are larger and more numerous in comparison to those of R. seeberi, and C. immitis can be cultured whereas R. seeberi cannot.10

Spontaneous regression of rhinosporidial growths has been noted in animals and in humans but is rare; therefore, medical and/ or surgical intervention is necessary. Surgical removal and electrodesication are the treatments of choice. Adjuvant medical therapies, such as ketoconazole, amphotericin B, trimethoprim-sulfadiazine, and sodium stibogluconate have been tried, with varied success.11 Data on antimicrobial drug resistance in R. seeberi are lacking. The strains obtained from human and animal rhinosporidiosis have shown genetic variations that might explain the variation of responses to some drugs.11

The only drug that appears to have clinical promise in treatment is dapsone. One study suggested that prolonged use of dapsone (100 mg/day for several months) may help prevent disease recurrence.12 It arrests the maturation of sporangia and promotes fibrosis in the stroma when used as an adjunct to surgery. It could be expected that presurgical dapsone would minimize the hemorrhage by its promotion of fibrosis, as well as preventing the colonization and infection of new sites after the release of endospores from surgically traumatized lesions.11 Postoperative use of dapsone has been reported to minimize or prevent recurrence.13

CONCLUSIONS

Our patient, who had recurrent nasopharyngeal lesions, underwent multiple endoscopic surgeries presented with multiple disseminated cutaneous lesions possibly with hematogenous spread.

REFERENCES

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CASE STUDY

Woodworm Bites in a 52-Year-Old Man
Claudio Conforti, MD;1,2 Maria Silvestre, MD;1 Caterina Dianzani, MD, PhD1

A 52-year-old man was admitted for clinical counseling at the Campus Bio-Medico in Rome in the early days of August 2017. He presented with multiple erythematous-hemorrhagic papules localized on the abdomen (Figure 1), back, arms, and thighs (Figure 2). The wide eruption was characterized by an intense reddish color that disappeared with diascopy. The patient stated that the eruption had appeared overnight after helping his mother-in-law dispose of a piece of antique furniture from her country villa the previous afternoon. Unexpectedly for the patient, the eruption did not cause intense itching during the night (it became more severe upon waking the next morning), and it did not affect his wife. (SKINmed. 2018;16:67–68)

The clinical presentation was extremely clear: the saliva of Scleroderma, with its anticoagulant and anesthetic properties, had created the typical bleeding pattern on the body of our patient. The differential diagnosis was very difficult. We excluded an infection caused by Pyemotes herfsi, in which lesions are generally smaller and less symptomatic.1 In the differential diagnosis, we also included scabies and bedbug bites.

The patient was not febrile, even though he complained general malaise and asthenia. The lymph nodes were neither swollen nor painful. Blood tests were normal except for an increase in the range of neutrophils and eosinophils. Antinuclear antibodies and viral serology were negative. Histopathologic examination of a skin sample showed the presence of edema in the dermis, and perivascular infiltration of neutrophils with the occasional presence of some eosinophils.

We treated the patient with antihistamines and topical corticosteroids, and there was a marked improvement by day 4.

DISCUSSION

Scleroderma domesticum is an insect belonging to the order Hymenoptera, family Betilidi. This insect has a slender body and is approximately 3 to 4 mm in length (Figure 3), not so different from the body of an ant. Its larvae are the common woodworm seen in furniture, where it also matures. Scleroderma domesticum attacks and feeds on the larvae of Anobium punctatum, Hylotrupes bajulus, Lasioderma serricorne, Nicobium castaneum, and Stegobium panicum.2 In absence of the usual prey, females of S. domesticum can bite humans, usually several times, on the limbs, chest, and back. The maximum incidence of activity is from April to June, especially in countries with a temperate climate.3

Despite having a worldwide distribution, S. domesticum is more prevalent in temperate zones.4 This may explain its spread in areas of Europe such as Italy and Spain. Numerous cases of S. domesticum bites have been reported in patients aged between 20 and 70 years in Italy. Both sexes were affected. A very small percentage of cases was related to Scleroderma brevicornis.3

Because the natural habitat of S. domesticum is represented by wooden objects, such as chairs, tables, and antique furniture, those most at risk of infestation by Scleroderma are restorers and antiquarians; for this reason, Scleroderma infestation is considered to be linked to professional environments.

The clinical response of a person stung by S. domesticum can vary from small local reactions to larger local reactions that may involve late-phase responses from IgE-mediated sensitivity.5,6 Scleroderma domesticum belongs to the order Hymenoptera, and this explains why its stings can cause disseminated eruptions. In our patient, the clinical presentation was extremely clear: erythematous maculae, papules, and severe itching. Other dermatozoonoses can be evaluated in the differential diagnosis, for example stings caused by mites, but this diagnosis was excluded because the patient did not present the classic nocturnal itching and pathognomonic lesions of scabies.

Clinical manifestations disappeared after 4 days of topical treatments with corticosteroids and oral antihistamines.
CONCLUSIONS

Considering the sting of *S. domesticum* as a possible diagnosis is important because timely treatment allows complete remission within a few days of clinical manifestations that typically result in loss of working days if left untreated. *Scleroderma domesticus* infestation occurs mainly in carpenters, antiquarians, and restorers; for this reason, it can be considered an occupational disease. A higher risk for reactions to Hymenoptera venom is reported in men.

In the evaluation of the patient’s response to insect stings, it is necessary to define whether reactions are local or systemic, to distinguish this kind of reaction from allergic reactions. These elements must be taken into consideration to achieve an early diagnosis, which will lead to reduced clinical manifestations and an improved quality of life for the patient.

REFERENCES


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Multiple Grouped Scalp Nodules in a Middle-Aged Man: A Rare Case of Angiolymphoid Hyperplasia with Eosinophilia

Lina Husienzad, BS;1 Ben Friedman, MD;2 Sherry X. Yang, MD;2 Amber Tran, MSN, CRNP;2 Jason B. Lee, MD2

An otherwise healthy man in his 50s presented complaining of pruritic lesions on the left side of his scalp. The lesions had slowly been growing in size over the preceding 30 years. They would occasionally bleed, and this is what ultimately prompted him to seek medical advice. Physical examination revealed multiple aggregated and soft, flesh-colored nodules on the left posterior auricular area of the scalp (Figure 1). No appreciable clinical lymphadenopathy was identified on examination. A shave biopsy of one of the nodules was performed for diagnostic clarification. (SKINmed. 2018;16:71–72)

Histopathologic examination revealed a patchy, nodular proliferation of vascular structures of varying thickness (Figure 2). Many of the vessels demonstrated prominent hobnail endothelial cells. A diffuse and polymorphous interstitial inflammatory infiltrate consisting of lymphocytes, plasma cells, and eosinophils was also observed. Given these findings, the patient was diagnosed with angiolymphoid hyperplasia with eosinophilia (ALHE). The patient was interested in treatment, and he was referred for surgical excision.

DISCUSSION

ALHE is a rare vasoproliferative tumor, characterized by a proliferation of blood vessels in the dermis and subcutaneous tissue.1 ALHE has a predilection for the skin and subcutaneous tissue of the head and neck; however, it has also been reported to occur at other sites, including the trunk, extremities, penis, and even visceral organs such as the colon.2 It has a reported mean age of onset of approximately 30 years, although it may present between the ages of 20 and 50.3 It is most commonly encountered in Asian populations, and has a slight female predominance.3

Patients with ALHE clinically present with a single or multiple aggregated flesh-colored-to-purple papules or nodules, which may be painful or pruritic.3 There is often associated pulsation and spontaneous bleeding, due to the prominent vascularity of the lesions.3 The characteristic histopathologic findings in ALHE are a proliferation of vascular structures with prominent hobnail endothelial cells among a dense background inflammatory infiltrate containing eosinophils and lymphocytes.4 The pathophysiology underlying ALHE is poorly understood. Some have suggested that it may be a reactive process to trauma or infection, whereas many now classify it as an arteriovenous malformation or other vasoproliferative disorder.4 Although it is considered to be a benign entity, it may persist and prove difficult to resolve even with treatment.3

Clinically, ALHE can resemble Kimura disease, and both were previously thought to represent the same disease entity at different ends of a spectrum; however, there is some evidence to suggest that they are actually clinically and pathologically distinct.5 Kimura disease often presents as a systemic illness and consists of a large subcutaneous mass in the head and neck region with...
72 Multiple Grouped Scalp Nodules

SKINmed. 2018;16:71–72

Figure 1. Clinical image showing multiple grouped, flesh-pink–colored nodules on the left postauricular region of the scalp.

Figure 2. Histopathologic images of the excised lesions. (A) Dome-shaped nodule demonstrating a proliferation of thin- and thick-walled vessels along with a patchy interstitial inflammatory infiltrate (hematoxylin and eosin stain, original magnification ×25). (B) Dilated vascular structures lined by hobnail endothelial cells with a mixed surrounding inflammatory infiltrate consisting of lymphocytes, plasma cells, and prominent eosinophils (hematoxylin and eosin stain, original magnification ×200).

associated lymphadenopathy, eosinophilia, elevated immunoglobulin E levels, and renal disease. Our patient lacked any signs of systemic disease, which made ALHE the more likely diagnosis in his case. Other entities that can resemble ALHE clinically include angiosarcoma, epithelioid hemangioendothelioma, Kaposi sarcoma, and pyogenic granuloma. These entities can easily be distinguished on histology.

Standard surgical excision is considered the most effective treatment to date for ALHE, although disease may recur in up to one-third of patients. The high recurrence rate may reflect the difficulty of identifying the margins of this highly vascular lesion. Pulsed-dye laser, carbon dioxide laser, Mohs micrographic surgery, and topical immunomodulators are other modalities that have shown variable success in treating ALHE. Cosmetic outcomes vary depending on the size and extent of the lesions.

REFERENCES

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CASE STUDY
Glucose Monitoring Dermopathy
Warren R. Heymann, MD

A 75-year-old man had been monitoring his glucose using a blood glucose monitoring system at the same body site for at least 20 years (>7300 needlesticks). The asymptomatic skin lesion had been present for many years. He used the same site because it hurt less than the fingers and bled well. His medical history was remarkable for diabetes mellitus, hypertension, coronary artery disease, and a pacemaker. His medications included glipizide, metformin, carvedilol, furosemide, lisinopril, amlodipine, clopidogrel, and aspirin. Physical examination revealed a brown, slightly raised, rough plaque with focal, punctate hemorrhagic crusts, on the distal area of the right thigh (Figure 1). The clinical differential diagnosis was more likely to be lichen simplex chronicus than pigmented Bowen's disease. A skin biopsy demonstrated an acanthotic epidermis with coarse collagen bundles in a thickened papillary dermis with extravasated erythrocytes, consistent with a dermal reparative reaction (Figure 2). (SKINmed. 2017;15:74–75)

Cutaneous manifestations of diabetes mellitus are well known to dermatologists and include necrobiosis lipoidica, diabetic dermopathy, diabetic bullae, scleredema diabeticorum, acanthosis nigricans, generalized granuloma annulare, eruptive xanthomatosis, and acquired perforating disorders. Glucose monitoring is essential in managing diabetes mellitus. A novel skin lesion secondary to repeated needlestick monitoring is reported.

DISCUSSION
Needlestick glucose monitoring has allowed for improved control of diabetes mellitus. Complications include pain at the injection site or infections such as Pseudomonas aeruginosa or Staphylococcus aureus. Hepatitis B and C may occur when needles are shared—a practice that should never happen. Injecting the needlestick into the fingertip yields the most accurate reading of serum glucose, although other body sites may be used at times when the glucose is likely not to rise or fall precipitously. Some manufacturers recommend rotating the site of testing.

CONCLUSIONS
Repeated needlestick injections at a solitary site for glucose monitoring may result in a hyperpigmented purpuric scar that could be mistaken for another neoplastic or inflammatory process. The purpuric component may be more prominent in patients who are anticoagulated. As there were no worrisome histologic features in our patient’s case, in the context of a precise history, reassurance is all that is necessary. It may be prudent to advise rotating the site of needlestick injection to avoid this complication of glucose monitoring.
Glucose Monitoring Dermopathy

ACKNOWLEDGMENT

Lawrence David Hall, MD, provided the histopathology photomicrograph.

REFERENCES


Figure 2. The histology of glucose-monitoring dermopathy, demonstrating an acanthotic epidermis with coarse collagen bundles in a thickened papillary dermis with extravasated erythrocytes, consistent with a dermal reparative reaction (hematoxylin and eosin stain, magnification ×200).

HISTORICAL DIAGNOSIS AND TREATMENT: EPITHELIOMA (Continued from page 53)

SYphilis secundaria

Syphilis pustulosa.

The general pustular syphilid is of much less common occurrence than the papular, and is observed most often in poorly nourished, debilitated, or anemic individuals, usually in the first three to eight months of the disease. Like the papular rash it may be the first recognized cutaneous manifestation; more often it follows or develops from a prior papular eruption. Pustules do not necessarily arise from papules, they may originate as pustules. When they appear in profusion their evolution is often preceded and accompanied by fever, headache and malaise. In the earlier rashes the pus is situated between the epidermis and the true skin. The later and relapsing forms are usually more limited in their distribution, the lesions are generally larger and the destruction deeper. In either case the pustules dry quickly with the formation of yellow, brown or greenish crusts which fall off and uncover characteristic dark-colored papules with a collarette of shredded epidermis at the periphery, or disclose shallow, punched out ulcers which may leave permanent scars when they heal. The pustular rash has many characteristics in common with the popular, namely, the color, the general distribution, the moderate grouping of the lesions, the tendency for new lesions to appear for several days or weeks and for the efflorescences to be nearly all of one size, nutilaria, small or large. In the milia-like pustular syphilid the lesions are usually profuse and situated about the hair follicles. A few large papules or pustules are nearly always present. The appearance of such an eruption is quite characteristic and not easily mistaken. Slightly larger pustules may resemble those of acne or varicella. The rash causes no subjective symptoms except that in the negro itching is sometimes complained of. DIAGNOSIS: The pustular syphilid may simulate any of the pustular non-specific skin lesions. Of the very greatest importance in diagnosis is the presence of other symptoms of syphilis. Points to be considered in differentiating the pustular syphilid from acne are, that acne is very chronic and localized upon the face and upper trunk, it rarely arises after thirty, comedones are usually plentiful, and the nodes may be deep and are of a dusky and not a coppery red. Feltm is distinguished by the sudden onset and intensity of its general symptoms, its acute course and definite duration, the shot-like feel of its papules and the full globular character of the pustules, the uniformity in character and development of the lesions and their predilection for the face, wrists and hands. In the exceedingly rare cases in which these points do not suffice for a differential diagnosis a serodiagnostic test will resolve all difficulties. TREATMENT: The general mercurial treatment is as a rule all that is required, though it is particularly in these cases that mercury vapor baths are serviceable.
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CASE STUDY

Bullosis Diabeticorum: A Neglected Bullous Dermatosis

Ramya Vangipuram, MD; Tiffany Hinojosa, MD; Daniel J. Lewis, BA; Christopher Downing, MD; Caleb Hixson, DO; Julio César Salas-Alanis, MD, PhD; Stephen K. Tyring, MD, PhD

A 75-year-old African-American man presented with a 3-year history of painless, fluid-filled blisters, for which his primary care physician had treated him with doxycycline, cephalixin, and topical corticosteroids, with no significant improvement. The blisters had ruptured spontaneously and healed with scarring. He denied antecedent trauma. His medical history was remarkable for insulin-dependent type 2 diabetes mellitus, hypertension, hypercholesterolemia, primary cutaneous melanoma status-post excision, and breast cancer status-post mastectomy and chemotherapy. Physical examination revealed nontender bullae, measuring up to 4 cm × 3 cm and containing serous fluid, on the anterior portion of both tibias (Figure 1). The Nikolsky sign was negative. There was no evidence of surrounding inflammation. A biopsy revealed subepidermal bullae formation with sparse inflammatory infiltrate (Figure 2). Direct and indirect immunofluorescence studies were negative for immunoglobulin (Ig) G, IgA, IgM, complement C3, C5b-9, and fibrinogen deposition. Culture of the bullous fluid was negative. (SKINmed. 2018;16:77–79)

Given these negative findings, the patient was diagnosed with bullous dermatosis (BD) and given topical mupirocin cream to prevent infection after the bullae ruptured. After 1 month, his condition had not changed.

DISCUSSION

BD is a rare, underreported cutaneous manifestation of diabetes mellitus.1,2 The disease is characterized by the spontaneous appearance of flaccid or tense bullae located on acral sites, often draining serous fluid.3 It is a relapsing and remitting disease. The bullae classically heal without scarring; our patient had an atypical presentation including healing with scarring. The microscopic location of the intercellular split varies and can occur in an intraepidermal or subepidermal manner.4 The pathogenesis of BD remains uncertain but may involve a response to trauma on acral locations as a complication of diabetic neuropathy and microangiopathy.2,3 Treatment of BD is typically supportive.1–4 The bullae should be kept intact to prevent secondary infection. Topical therapy is generally not required, although topical antibiotics such as mupirocin may prevent infection. In cases of tissue necrosis, debridement and tissue grafting should be considered.1 Reports of osteomyelitis and infection requiring amputation arising at the site of bullous disease illustrate the importance of monitoring for secondary infection and providing aggressive wound healing intervention, as with diabetic ulcers, if the bullae become unroofed.2

CONCLUSIONS

The clinical features of BD are nonspecific. The bullae seen in bullous pemphigoid, epidermolysis bullosa acquista, and porphyria cutanea tarda may mimic those of BD (Table). Microscopy and direct and indirect immunofluorescence features are used to exclude these diseases. In our case, the biopsy findings were consistent with diabetic bullae and supported by the history of diabetes. BD is a
Figure 1. Tense, fluid-filled bulla on the lower extremity.

Figure 2. Biopsy showing subepidermal bulla formation and sparse inflammatory infiltrate (hematoxylin and eosin stain, original magnification ×20).

Table. Differential Diagnosis of Bullous Diabeticorum

<table>
<thead>
<tr>
<th>Dermatosis</th>
<th>Histology Findings</th>
<th>DIF</th>
<th>IDIF</th>
<th>Clinical Features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pemphigus vulgaris5</td>
<td>Intraepidermal bullae, eosinophils, loss of</td>
<td>Reticular pattern of IgG and</td>
<td>IgG binds to desmoglein</td>
<td>Presents in skin and oral mucosa; positive Nikolsky sign; flaccid bullae that</td>
</tr>
<tr>
<td></td>
<td>intercellular attachments in basement membrane</td>
<td>complement deposits around</td>
<td></td>
<td>rupture easily leaving shallow erosions</td>
</tr>
<tr>
<td>Bullous pemphigoid1</td>
<td>Subepidermal bullae, eosinophil-predominant</td>
<td>Linear-pattern (N-serrated)</td>
<td>IgG binds to hemidesmosomes</td>
<td>Presents only in skin; negative Nikolsky sign; tense bullae that do not rupture</td>
</tr>
<tr>
<td></td>
<td>inflammatory infiltrate</td>
<td>of IgG and complement C3</td>
<td>and upper lamina lucida</td>
<td>easily</td>
</tr>
<tr>
<td>Dermatitis herpetiformis6</td>
<td>Subepidermal bullae, neutrophilic microabscesses</td>
<td>Granular pattern of IgA</td>
<td></td>
<td>Pruritic, grouped vesicles and bullae</td>
</tr>
<tr>
<td>Epidermolysis bullosa</td>
<td>Subepidermal bullae, mixed inflammatory cell</td>
<td>Linear pattern (U-serrated)</td>
<td>IgG binds to dermal floor</td>
<td>Mild mucosal involvement; healing with dense scars at trauma areas</td>
</tr>
<tr>
<td>acquisita7</td>
<td>infiltrate</td>
<td>of IgG, IgM or IgA, and</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>C3 deposition at basement</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>membrane</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Porphyria cutanea tarda8</td>
<td>Free-cell subepidermal bullae, festooning of the</td>
<td>Granular pattern of IgG,</td>
<td>—</td>
<td>Sunlight exposure with subsequent appearance of lesions; additional hair growth</td>
</tr>
<tr>
<td></td>
<td>dermal papillae</td>
<td>and C3 deposition at the</td>
<td></td>
<td>and pigmentation changes</td>
</tr>
<tr>
<td>Drug-induced linear IgA9</td>
<td>Subepidermal bullae, neutrophilic microabscesses</td>
<td>Linear pattern of IgA at</td>
<td>IgA binds to basement membrane</td>
<td>Transient pruritus or burning preceding lesions; commonly precipitated by sulfa</td>
</tr>
<tr>
<td></td>
<td></td>
<td>basement membrane</td>
<td></td>
<td>drugs, β-lactams, and phenytoin</td>
</tr>
<tr>
<td>Bullosis diabeticorum</td>
<td>Intraepidermal or subepidermal bullae, sparse</td>
<td>Negative for IgG, IgA, IgM,</td>
<td>Negative for IgG, IgA, IgM,</td>
<td>Spontaneous appearance of tense or flaccid bullae on acral sites; often drain</td>
</tr>
<tr>
<td></td>
<td>inflammatory infiltrate</td>
<td>C3, C5b-9, and fibrinogen</td>
<td>C3, C5b-9, and fibrinogen</td>
<td>serous fluid; self-remitting</td>
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<td></td>
<td></td>
<td>deposition</td>
<td>deposition</td>
<td></td>
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</table>

Abbreviations: C, complement; DIF, direct immunofluorescence; IDIF, indirect immunofluorescence; Ig, immunoglobulin.
diagnosis of exclusion. A detailed study to differentiate the histopathology of other BDs may help in understanding the evolution of this condition, resulting in improved management and prevention.

REFERENCES

1 Lipsky BA, Baker PD, Ahroni JH. Diabetic bullae: 12 cases of a purportedly rare cutaneous disorder. *Int J Derma


8 Schulenburg-Brand D, Katugampola R, Anstey AV, Badmin

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CONTRAINdications
The LEVULAN KERASTICK Topical Solution plus blue light illumination using the BLU-U Blue Light Photodynamic Therapy Illuminator is contraindicated in patients with cutaneous photosensitivity at wavelengths of 400-450 nm, porphyria or known allergies to porphyrins, and in patients with known sensitivity to any of the components of the LEVULAN KERASTICK for Topical Solution.

WARNINGS AND PRECAUTIONS
Photosensitivity
During the time period between the application of LEVULAN KERASTICK Topical Solution and exposure to activating light from the BLU-U Blue Light Photodynamic Therapy Illuminator, the treatment site will become photosensitive. After LEVULAN KERASTICK Topical Solution application, patients should avoid exposure of the photosensitive treatment sites to sunlight or bright indoor light (e.g., examination lamps, operating room lamps, tanning beds, or lights at close proximity) during the period prior to blue light treatment. Exposure may result in a stinging and/or burning sensation and may cause erythema and/or edema of the lesions. Before exposure to sunlight, patients should, therefore, protect treated lesions from the sun by wearing a wide-brimmed hat or similar head covering of light-opaque sunscreen. Sunscreens will not protect against photosensitivity reactions caused by visible light. It has not been determined if perspiration can spread the LEVULAN KERASTICK Topical Solution outside the treatment site to eye or surrounding skin.

Application of LEVULAN KERASTICK Topical Solution to perilesional areas of photodamaged skin of the face or scalp may result in photosensitization. Upon exposure to activating light from the BLU-U Blue Light Photodynamic Therapy Illuminator, such photosensitized skin may produce a stinging and/or burning sensation and may become erythematous and/or edematous in a manner similar to that of actinic keratoses treated with LEVULAN KERASTICK Photodynamic Therapy. Because of the potential for skin to become photosensitized, the LEVULAN KERASTICK should be used by a qualified health professional to apply drug only to actinic keratoses and not perilesional skin. If for any reason the patient cannot return for blue light treatment during the prescribed period after application of LEVULAN KERASTICK Topical Solution (14 to 18 hours), the patient should call the doctor immediately. The patient should also continue to avoid exposure of the photosensitized lesions to sunlight or prolonged or intense light for at least 40 hours. If stinging and/or burning is noted, exposure to light should be reduced.

Irritation
The LEVULAN KERASTICK Topical Solution contains alcohol and is intended for topical use only. Do not apply to the eyes or to mucous membranes. Excessive irritation may be experienced if this product is applied under occlusion.

Coagulation Defects
The LEVULAN KERASTICK for Topical Solution has not been tested on patients with inherited or acquired coagulation defects.

ADVERSE REACTIONS
In Phase 3 studies, no non-cutaneous adverse events were found to be consistently associated with LEVULAN KERASTICK Topical Solution application followed by blue light exposure.

Photodynamic Therapy Response: The constellation of transient local symptoms of stinging and/or burning, itching, erythema and edema as a result of LEVULAN KERASTICK Topical Solution plus BLU-U treatment was observed in all clinical studies of LEVULAN KERASTICK for Topical Solution Photodynamic Therapy for actinic keratoses treatment. Stinging and/or burning subsided between 1 minute and 24 hours after the BLU-U Blue Light Photodynamic Therapy Illuminator was turned off, and appeared qualitatively similar to that perceived by patients with erythropoietic protoporphyria upon exposure to sunlight. There was no clear drug dose or light dose dependent change in the incidence or severity of stinging and/or burning.

In two Phase 3 trials, the sensation of stinging and/or burning appeared to reach a plateau at 6 minutes into the treatment. Severe stinging and/or burning at one or more lesions being treated was reported by at least 50% of the patients at some time during treatment. The majority of patients reported that all lesions treated exhibited at least slight stinging and/or burning. Less than 3% of patients discontinued light treatment due to stinging and/or burning.

In the Phase 3 trials, the most common changes in lesion appearance after LEVULAN KERASTICK for Topical Solution Photodynamic Therapy were erythema and edema. In 96% of active treatment patients, some or all lesions were erythematous shortly after treatment, while in 79% of vehicle treatment patients, some or all lesions were erythematous. In 35% of active treatment patients, some or all lesions were edematous, while no vehicle-treated patients had edematous lesions. Both erythema and edema resolved to baseline or improved by 4 weeks after therapy. LEVULAN KERASTICK Topical Solution application to photodamaged perilesional skin resulted in photosensitization of photodamaged skin and in a photodynamic response (see Warnings and Precautions).

Other Localized Cutaneous Adverse Experiences: Table 1 depicts the incidence and severity of cutaneous adverse events in Phase 3 studies, stratified by anatomic site treated.

<table>
<thead>
<tr>
<th>Site</th>
<th>Treatment</th>
<th>Incidence</th>
<th>Severity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Face</td>
<td>Treatment</td>
<td>Incidence</td>
<td>Severity</td>
</tr>
<tr>
<td>Face</td>
<td>Control</td>
<td>Incidence</td>
<td>Severity</td>
</tr>
</tbody>
</table>

Adverse Experiences Reported by Body System:
In the Phase 3 studies, 7 patients experienced a serious adverse event. All were deemed remote or not related to treatment. No clinically significant patterns of clinical laboratory changes were observed for standard serum chemical or hematologic parameters in any of the controlled clinical trials.

OVERDOSAGE
LEVULAN KERASTICK Topical Solution Overdose LEVULAN KERASTICK Topical Solution overdose has not been reported. In the unlikely event that the drug is ingested, monitoring and supportive care are recommended. The patient should be advised to avoid accidental exposure to intense light sources for at least 40 hours after ingestion. The consequences of exceeding the recommended topical dosage are unknown.

BLUE Light Overdose
There is no information on overdose of blue light from the BLU-U Blue Light Photodynamic Therapy illuminator following LEVULAN KERASTICK Topical Solution application.

Information for Patients
LEVULAN KERASTICK Photodynamic Therapy for Actinic Keratoses
- The first step in LEVULAN KERASTICK Photodynamic Therapy (PDT) for actinic keratoses is application of the LEVULAN KERASTICK Topical Solution to actinic keratoses located on the patient’s face or scalp.
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