Dermoscopy in family medicine: A primer

Dermoscopy allows you to see deeper into the skin than with the naked eye. Here’s how you can make use of it to spot malignant conditions sooner.

Dermoscopy, the use of a handheld instrument to magnify the skin 10-fold while providing a light source, is a quick, useful, cost-effective tool for detecting melanoma in family medicine.1-4

The device, which allows the physician to visualize structures below the stratum corneum that are not routinely discernible with the naked eye, can be attached to a smartphone so that photos can be taken and reviewed with the patient. The photo can also be reviewed after a biopsy result is obtained.

Its use among non-dermatologist US physicians appears to be relatively low, but rising. One small study of physicians working in family medicine, internal medicine, and plastic surgery found that only 15% had ever used a dermatoscope and 6% were currently using one.5

As a family physician, you can expand your diagnostic abilities in dermatology with the acquisition of a dermatoscope (FIGURE 1) and some time invested in learning to interpret visible patterns. With that in mind, this review focuses on the diagnosis of skin cancers and benign growths using dermoscopy. We begin with a brief look at the research on dermoscopy and how it is performed. From there, we’ll detail an algorithm to guide dermoscopic analysis. And to round things out, we provide guidance that will help you to get started. (See “Choosing a dermatoscope—and making the most of it,” page E3 and “To learn more about dermoscopy ...” page E7.)

What the research says
Dermoscopy improves sensitivity for detecting melanoma over the naked eye alone; it also allows for the detection of melanoma at earlier stages, which improves prognosis.6

A meta-analysis of dermoscopy use in clinical settings showed that, following training, dermoscopy increases the average sensitivity of melanoma diagnosis from 71% to more than 90% without a significant decrease in specificity.7 In a study of 74 primary care physicians, there was an improvement in both clinical and dermo-

FIGURE 1
A sampling of dermatoscopes

Dermatoscopes come in all shapes and sizes, with polarized and/or nonpolarized light sources. Some can also be attached to a camera or smartphone.
The exam begins by choosing 1 of 3 modes of dermoscopy

Dermatoscopes can have a polarized or non-polarized light source. Some dermatoscopes combine both types of light (hybrid dermatoscopes; see “Choosing a dermatoscope—and making the most of it,” page E3.)

- nonpolarized contact dermoscopy
- polarized contact dermoscopy
- polarized non-contact dermoscopy.

Dermatoscopes with nonpolarized light require direct skin contact and a liquid interface (eg, alcohol, gel, mineral oil) between the scope’s glass plate and the skin for the visualization of subsurface structures. In contrast, dermatoscopes with polarized light do not require direct skin contact or a liquid interface; however, contacting the skin and using a liquid interface will provide a sharper image.

Two major algorithms guide dermoscopic analysis

The first of 2 major algorithms that can be used to guide dermoscopic analysis is a modified pattern analysis put forth by Kittler. This descriptive system based on geometric elements, patterns, colors, and clues guides the observer to a specific diagnosis without categorizing lesions as being either melanocytic or nonmelanocytic. Because this is not the preferred method of the authors, we will move on to Method 2.

The second method, a 2-step algorithm, is a qualitative system that guides the observer through differentiating melanocytic from nonmelanocytic lesions in order to differentiate nevi from melanoma (FIGURE 2). At the same time, it serves as an aid to correctly diagnose non-melanocytic lesions. The 2-step algorithm forms the foundation for the dermoscopic evaluation of skin lesions in this article.
Not all expert dermoscopists employ structured analytical systems or methods to reach a diagnosis. Because of their vast experience, many rely purely on pattern recognition. But algorithms can facilitate non-experts in dermoscopy in the differentiation of nevi from melanoma or, simply, in differentiating the benign from the malignant.

Although each algorithm has its unique criteria, all of them require training and practice and familiarity with the terms used to describe morphologic structures. The International Dermoscopy Society recently published a consensus paper designating some terms as preferred over others.11

Step 1: Melanocytic vs non-melanocytic
Step 1 of the 2-step algorithm requires the observer to determine whether the lesion is melanocytic (ie, originates from melanocytes and, therefore, could be a melanoma) or non-melanocytic in origin.

A melanocytic lesion usually will display at least 1 of the following structures:
- pigment network (FIGURE 3A) (This can include angulated lines.)
- negative network (FIGURE 3B) (hypopigmented lines connecting pigmented structures in a serpiginous fashion)
- streaks (FIGURE 3C)
- homogeneous blue pigmentation (FIGURE 3D)
- globules (aggregated or as a peripheral rim) (FIGURE 3E)
- pseudonetwork (facial skin) (FIGURE 3F)
- parallel pigment pattern (acral lesions) (FIGURE 3G).

Exceptions. Sometimes, nonmelanocytic lesions will present with pigment network. Dermatofibromas, for example, are one exception in which the pattern trumps the network. Two other exceptions are solar lentigo and supernumerary or accessory nipple.

If the lesion does not display any structure, it is considered structureless. In these cases, proceed to the second step to rule out a melanoma.

Choosing a dermatoscope—and making the most of it

1. Consider acquiring a hybrid dermatoscope. Nonpolarized dermatoscopes (NPDs) and polarized dermatoscopes (PDs) provide different but complementary information. PDs enable users to identify features such as vessels and shiny white structures that are highly indicative of skin cancer. Because PDs are highly sensitive for detecting skin cancer and do not require a liquid interface or direct skin contact, they are the ideal dermatoscopes to use for skin cancer screening.

However, maintaining the highest specificity requires the complementary use of NPDs, which are better at identifying surface structures seen in seborrheic keratoses and other benign lesions. Thus, if the aim is to maintain the highest diagnostic accuracy for all types of lesions, then the preferred dermatoscope is a hybrid that permits the user to toggle between polarized and nonpolarized features in one device.

2. Choose a dermatoscope that attaches to your smartphone and/or camera. This helps you capture digital dermoscopic images that can be analyzed on a larger screen, which permits:
- enlarging certain areas for in-depth analysis of structures and patterns
- sharing the image with the patient to explain why a biopsy is, or isn’t, needed
- sharing the image with a colleague for the purpose of a consult or a referral, or using the images for teaching purposes
- saving the images in order to follow lesions over time when monitoring is indicated
- ongoing learning. After each biopsy result comes back, we recommend correlating the dermoscopic images with the biopsy report. If your suspected diagnosis was correct, this reinforces your knowledge. If the pathology diagnosis is unexpected, you can learn by revisiting the original images to look for structures or patterns you may have missed upon first examination. You may even question the pathology report based on the dermoscopy, prompting a call to the pathologist.
- keeping a safe distance from the patient when looking for scabies mites.
FIGURE 3
Features of melanocytic lesions

Melanocytic lesions may display pigment network (A), negative network (B), streaks, such as around the perimeter of this Spitz nevus (C), homogeneous blue pigmentation (D), globules (aggregated or as a peripheral rim; E), pseudonetwork (facial skin; F), or a parallel pigment pattern (acral lesions; G).

PHOTOS COURTESY OF: RICHARD P. USATINE, MD, AND ASHFAQ A. MARGHOOB, MD
Doesn’t meet criteria for a melanocytic lesion?
If the lesion does not reveal any of the criteria for a melanocytic lesion, then look for structures seen in nonmelanocytic lesions: dermatofibromas; seborrheic keratosis; angiomomas and angiokeatromas; sebaceous hyperplasia; clear-cell acanthomas; basal cell carcinomas (BCCs); and squamous cell carcinomas (SCCs).

Benign nonmelanocytic lesions
Dermatofibromas are benign symmetric lesions that feel firm and may dimple upon application of lateral pressure. They are fibrotic scar-like lesions that present with 1 or more of the following dermoscopic features (FIGURE 4):

- peripheral pigment network, due to increased melanin in keratinocytes
- homogeneous brown pigmented areas
- central scar-like area
- shiny white lines
- vascular structures (ie, dotted, polymorphous vessels), usually seen within the scar-like area
- ring-like globules, usually seen in the zone between the scar-like depigmentation and the peripheral network. They correspond to widened hyperpigmented rete ridges.

Seborrheic keratosis (SK) is a benign skin growth that often has a stuck-on appearance (FIGURE 5). Features often include:

- multiple (>2) milia-like cysts
- comedo-like openings
- a network-like structure that corresponds to gyri and sulci and which in some cases can create a cerebriform pattern
- fingerprint-like structures
- moth-eaten borders
- jelly sign. This consists of semicircular u-shaped structures that have a smudged appearance and are aligned in the same direction. The appearance resembles jelly as it is spread on a piece of bread.
- hairpin (looped or twisted-looped) vessels surrounded by a white halo.

CONTINUED
Other clues include a sharp demarcation and a negative wobble sign (which we’ll describe in a moment). The presence or absence of a wobble sign is determined by using a dermatoscope that touches the skin. Mild vertical pressure is applied to the lesion while moving the scope back and forth horizontally. If the lesion slides across the skin surface, the diagnosis of an epidermal keratinocytic tumor (ie, SK) is favored. If, on the other hand, the lesion wobbles (rolls back and forth), then the diagnosis of a neoplasm with a dermal component (ie, intradermal or compound nevus) is more likely.

Angiomas and angiokeratomas. Angiomas demonstrate lacunae that are often separated by septae (FIGURE 6). Lacunae can vary in size and color. They can be red, red-white, red-blue, maroon, blue, blue-black, or even black (when thrombosis is present).

Angiokeratomas (FIGURE 7) can reveal lacunae of varying colors including black, red, purple, and maroon. In addition, a blue-whitish veil, erythema, and hemorrhagic crusts can be present.

Sebaceous hyperplasia is the overgrowth of sebaceous glands. It can mimic BCC on the face. Sebaceous hyperplasia presents with multiple vessels in a crown-like arrangement that do not cross the center of the lesion. The sebaceous glands resemble popcorn (FIGURE 8).

Clear-cell acanthoma is a benign erythematous epidermal tumor usually found on the leg with a string-of-pearls pattern. This pattern is vascular so the pearls are red in color (FIGURE 9).

Malignant nonmelanocytic lesions
BCC is the most common type of skin cancer. Features often include:

- spoke-wheel-like structures or concentric structures (FIGURE 10A)
- leaf-like areas (FIGURE 10B)
- arborizing vessels (FIGURE 10B AND 10C)
- large blue-gray ovoid nest (FIGURE 10A)
- multiple blue-gray non-aggregated globules
- ulceration or multiple small erosions
- shiny white structures and strands (FIGURE 10C).

Additional dermoscopic clues include
short, fine, superficial telangiectasias and multiple in-focus dots in a buck-shot scatter distribution.

**Squamous cell carcinomas (SCCs)** of the skin are keratinizing malignant tumors. Each SCC generally has some of the following features (FIGURE 11):

- dotted and/or glomerular vessels,
FIGURE 12

Benign nevi patterns

- Diffuse reticular
- Patchy reticular
- Peripheral reticular with central hypopigmentation
- Peripheral reticular with central hyperpigmentation

*Benign patterns encountered in many acquired nevi and dysplastic nevi. Blue nevi, some Spitz nevi, and congenital melanocytic nevi can also manifest some of these patterns.

**Nevi with this pattern should be interpreted with caution.

FIGURE 13

Melanoma-specific structures

- Atypical network
- Streaks (pseudopods and radial streaming)
- Negative pigment network
- Shiny white lines (crystalline structures)
- Atypical dots and/or globules
- Off-centered blotch
- Peripheral tan structureless areas
- Blue-white veil overlying raised areas
- Regression structures
  - Blue-white veil overlying macular areas; scar-like areas and/or peppering
- Atypical vascular structures
  - Dotted vessels, serpentine vessels, polymorphous vessels, milky-red areas, red globules, corkscrew vessels
- Polygonal structures (zig-zag lines)

Commonly distributed focally at the periphery. They can also be diffuse or aligned linearly within the lesion.

- Scale (yellow or white)
- Rosettes (seen with polarized light)
- White circles or keratin pearls
- Brown circles
- Ulcerations
- Brown dots or globules arranged in a linear configuration.

Step 2: It’s melanocytic, but is it a nevus or a melanoma?

If, by following Step 1 of the algorithm, the lesion is determined to be of melanocytic origin, then one proceeds to Step 2 to decide whether the growth is a nevus, a suspicious lesion, or a melanoma. For this purpose, several additional algorithms are available.\(^\text{12-17}\)

- **Benign nevi** tend to manifest with 1 of the following 10 patterns: (FIGURE 12)
  1. Diffuse reticular
  2. Patchy reticular
  3. Peripheral reticular with central hypopigmentation
FIGURE 14
Melanomas with melanoma-specific structures

Each of these melanomas demonstrates at least 2 melanoma-specific structures. The first (A) shows an atypical network with peripheral atypical dots and globules; (B) shows a structureless area and negative network, along with a blue-white veil regression structure, among other features; (C) demonstrates streaks, an atypical network, shiny white lines, and peppering; (D) reveals shiny white lines and a peripheral tan structureless area; (E) shows polygonal lines and scar-like areas, among other features; (F) shows regression structures, peppering, shiny white lines, and atypical dots and globules; (G) features streaks, a peripheral tan structureless area, and milky red areas; and (H) reveals irregularly distributed atypical globules, off-centered blotches, a scar-like area, and streaks.

4. peripheral reticulate with central hyperpigmentation
5. homogeneous
6. peripheral globules/starburst. It has been suggested that lesions that show starburst morphology on dermoscopy require complete excision and follow-up since 13% of Spitzoid-looking symmetric lesions in patients older than 12 years were found to be melanoma in one study.  
7. peripheral reticulate with central globules
8. globular
9. 2-component
10. symmetric multicomponent (this pattern should be interpreted with caution, and a biopsy is probably warranted for dermoscopic novices).
Melanomas tend to deviate from the benign patterns described earlier. Structures in melanomas are often distributed in an asymmetric fashion (which is the basis for diagnosis in many of the other algorithms), and most of them will reveal 1 or more of the melanoma-specific structures (FIGURE 13). The melanomas in FIGURES 14 A-H each show at least 2 melanoma-specific structures. On the face or sun-damaged skin, melanoma may present with grey color, a circle-in-circle pattern, and/or polygonal lines (FIGURE 15). Note that melanoma on the soles or palms may present with a parallel ridge pattern (FIGURE 16).

How to proceed after the evaluation of melanocytic lesions
After evaluating the lesion for benign patterns and melanoma-specific structures, there are 3 possible pathways:

1. The lesion adheres to one of the nevi patterns and does not display a melanoma-specific structure. You can reassure the patient that the lesion is benign.
2. The lesion:
   A. Adheres to one nevus pattern, but also displays a melanoma-specific structure.
   B. Does not adhere to any of the benign patterns and does not have any melanoma-specific structures.

   This is considered a suspicious lesion, and the choices of action include performing a biopsy or short-term monitoring by comparing dermoscopic images over a 3-month interval. (Caveat: Never monitor raised lesions because nodular melanomas can grow quickly and develop a worsened prognosis in a short time. Instead you’ll want to biopsy the lesion that day or very soon thereafter.)
3. The lesion deviates from the benign patterns and has at least 1 melanoma-specific structure. Biopsy the lesion to rule out melanoma.

A bonus: Diagnosing scabies
Increasingly, dermoscopy is being used in the diagnosis of many other skin, nail, and hair problems. In fact, one great bonus to owning a dermatoscope is the accurate diagnosis of scabies. Dermoscopy can be helpful
in detecting the scabies mite without having to scrape and use the microscope. Moreover, the sensitivity and specificity of a dermoscopic diagnosis is higher than for scraping and microscopy.19

What you’ll see

The anterior legs and mouth parts of the mite resemble a triangle (arrowhead, delta-wing jet) (FIGURE 17). Look for a burrow, and the mite can be seen at the end of the burrow as a faint circle with a leading darker triangle. The burrow itself has a distinctive pattern that has more morphology than an excoriation and has been described as the contrail of a jet plane. Using a dermatoscope attached to your smartphone allows you to magnify the image even further while maintaining a safe distance from the mite.

2 scabies mites and burrows

Two scabies mites and their burrows as seen with dermoscopy. Burrows often resemble the contrails of a jet plane. The dark triangles or chevrons at one end of the burrows represent the mite’s head and front legs.

References


CORRESPONDENCE

Richard P. Usatine, MD, 903 W. Martin, Skin Clinic – Historic Building, San Antonio, TX 78207; usatine@uthscsa.edu.

FIGURE 17