Localized Cutaneous Argyria: A Report of 2 Cases

Argyria cutánea localizada. A propósito de 2 casos

To the Editor:

Argyria is a rare disease caused by silver deposition in the body. Most reports of occupational argyria in the literature describe the generalized form. We report on 2 patients who developed localized argyria after more than 30 years of contact with silver. The clinical appearance of these lesions requires us to make a differential diagnosis including deep melanocytic lesions, especially melanoma metastases.

The first patient was a 75-year-old man with no previous history of interest who presented with asymptomatic blue macules that had begun to appear 5-10 years previously on the dorsum of the fourth finger of the right hand and on the palmar surface of the fifth finger of the right hand (Fig. 1, A and B). Dermoscopy revealed lesions with a homogeneous blue pattern (Fig. 1C). The second patient was a 70-year-old man who also presented with blue-gray macules on the dorsum of both hands that had begun to appear 5 years previously (Fig. 1D). Biopsy disclosed multiple dark brown strands and granules with a periadnexal distribution around the secretory coils of the sweat glands and an interstitial distribution with deposits along the elastic fibers of the dermis (Fig. 2, A, B, and C). Staining for melanocytes was negative. Dark-field microscopy revealed characteristic brilliant specks (Fig. 2D). After more exhaustive questioning, both patients stated that they had worked as jewelers for more than 30 years and habitually handled silver. They were diagnosed with localized cutaneous argyria and, after opting not to receive treatment, remain under follow-up. Two years after diagnosis, no changes have been detected in the lesions.

Argyria is an uncommon condition today, especially since silver-containing systemic medication, which was once used to treat a variety of conditions (including syphilis), is no longer prescribed. Nevertheless, a series of mechanisms has been described in recent years by which silver can unintentionally enter the body through the skin, including the use of earrings, body piercing, dental amalgam, topical silver salts, or acupuncture needles. Prolonged occupational exposure to silver occurs in individuals working in the manufacturing industry and photographic processing, as well as silversmiths and goldsmiths.

Clinically, argyria is classed as generalized or localized. Generalized argyria is produced by the ingestion or inhalation of silver compounds that subsequently reach internal organs and the skin, where they cause permanent diffuse blue-gray coloration in photoexposed areas.

Localized argyria is caused by direct external contact with silver; the most commonly affected areas are the hands, eyes, and the mucosas. The mechanism of penetration is unknown, although it is thought to be via the eccrine sweat ducts, since the metal is most concentrated around the secretory portion of the gland. The fact that accumulation of silver does not respond to chelators, as well as the relatively low frequency of systemic argyria in patients with intact skin who come into contact with silver, suggests that silver is deposited in the dermis in a chemically stable and apparently inert form. Physical examination reveals multiple asymptomatic blue-gray macules with a nonspecific homogeneous blue pattern on dermoscopy. Histology reveals apparently normal skin. However, increased magnification reveals multiple brown-gray strands and granules distributed between collagen fibers, within histocytes, in the basement membrane of the eccrine sweat glands, and along the elastic fibers of the dermis in a pattern resembling chains of streptococci (B and C) (hematoxylin-eosin, original magnification ×200). Dark-field microscopy shows the characteristic “starry sky” image (D).
In a detailed review article, Velasco et al. (J. Garcias-Ladaria) have focused on the significant impact on the healing process of glucocorticoid (GC) activity. Current research on new treatments is focusing on both the blockade of GC receptors and on the activity of 11β-hydroxysteroid dehydrogenase-1 (11β-HSD1), the enzyme responsible for the peripheral conversion of cortisone to its active form cortisol. The presence of high GC levels is associated with delayed cutaneous wound healing and barrier recovery after mechanical disruption. Using models based on exogenous GC administration or endogenous release secondary to psychological stress, several authors have demonstrated the inhibitory action of GCs on fibroblast proliferation. In both models, systemic GC blockade improved wound healing and barrier recovery.

Because of the potential complications of systemic blockade of GC activity, current research on new treatments is focusing on both the blockade of GC receptors and on the activity of 11β-hydroxysteroid dehydrogenase-1 (11β-HSD1), the enzyme responsible for the peripheral conversion of cortisone to its active form cortisol. Given the high levels of GC observed in diabetic patients and the recent suggestion of a possible association between GC activity and systemic complications in diabetes mellitus, topical treatments that block such activity would appear to be a promising therapeutic tool in wound healing. The dermatological application of such topical treatments could therefore be very useful in the treatment of leg ulcers. This is a field in which, as Velasco has pointed out, dermatologists should play a greater role.

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References


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