Agminated Dermal Melanocytosis in the Territory of Ota’s Nevus

Melanocitosis dérmica agminada en el territorio del nevus de Ota

To the Editor:

Together with Mongolian spot, Ota’s naevus, Ito’s naevus, blue nevi are believed to represent a dermal arrest in embryonal migration of neural crest melanocytes that fail to reach epidermis. They usually appear in childhood but sometimes later in life. After a trauma, damaged melanocytes and inflammatory cells can release alpha-Melanocyte-stimulating hormone, that could influence nevus cells to form larger nests and macroscopically visible nevi. Blue nevi usually are solitary bluish-pigmented neoplasms, rarely are multiple and clustered in a well-circumscribed area, as agminated subtype. The Latin term “agmen” means “army” and indicates this rare flat or raised arranged group of nevi. These lesions are distributed on trunk, extremities and head/neck. A twenty-year-old Bulgarian, phototye IV boy presented with multiple asymptomatic pigmented lesions on right hemiface involving the three major branches of trigeminal nerve, developed since 18 months of age after a trauma. He underwent surgical treatments to reduce the involved area. Family history for skin cancer was negative. We observed approximately 100 maculo-papular, blue and brown lesions of different sizes, forming a cluster of nevi (Fig. 1a). The skin around papules was not discolored nor indurated. The surgical scar was not free of pigmented process. Dermoscopy revealed homogeneous pattern with diffuse brownish areas, regular network at the periphery and numerous regularly distributed small dots or homogeneous blue-gray pattern (Fig. 1b). No background pigmentation was noted between lesions. Moreover we performed Reflectance Confocal Microscopy (RCM) and a single lesion was excised for histological evaluation, disclosing typical findings of blue nevi. RCM displayed a regular epidermal architecture with a typical honeycomb pattern. At the spinous- granular layer we observed bright and polygonal keratinocytes with dark central nuclei. At basal layer we observed a single layer of refractive cells with cobblestone pattern. The dermo-epidermal junction was preserved with dermal papillae surrounded by luminous cells, with round nucleous and elongated cells, correlating to inflammatory cells and melanophages. Within upper dermis, we noticed irregular bright area corresponding to collagen bundles. Irregular refractive structures with unvisible nucleous were observed within dermal papillae of reticular dermis corresponding melanophages, inflammatory cells and dendritic melanocytes (Fig. 1c). Histologic examination revealed deep and concentrated dermal fusiform or dendritic-like melanocytes, not arranged in clusters. Thickening of the connective tissue with melanophages was also observed. No bluish stain of the ipsilateral eye was observed and ophthalmic evaluation demonstrated a stromal corneal melanosis in the caruncle and corneal-limbal complex. Optico Coherence Tomography (OCT) visualized hyperreflective regular structure with definite margins in both regions (Fig. 1d). No pigmented areas were found under otolaryngological examination. Nuclear Magnetic Resonance displayed normal morphological and dimensional features and did not reveal signal intensity alterations of brain parenchyma. We report a case of a boy with congenital large agminated blue pigmented lesion within area of Ota’s nevus. The nevus enclosed distribution of the first, second divisions of the trigeminal nerve, but also extended into the third branch, which is rarely seen in Ota’s nevus. Physical exam revealed several maculo-papular blue and brown lesions, but the skin between papules appeared not discolorated nor indurated. No bluish discoloration of the eye was observed, but a stromal corneal melanosis in the caruncle and corneal-limbal complex is also present. Furthermore no oral mucosal or neurological lesions were noted. Moreover, histologic examination showed dermal melanocytes are deeply located and densely packed. These features were consistent with diagnosis of blue nevus. However, it would appear that the blue nevus, Mongolian spot and the nevus of Ota are closely related and possibly represent different entities within the spectrum of blue lesions. Considering these results and treatment limitations due lesion size, a 3-months-follow-up of patient was followed, carrying out multidisciplinary management to rule out malignant changes. Dermoscopy is important for early detection of malignant transformation: agminated blue nevi are commonly characterized by homogeneous pattern with diffuse brownish areas, in combination with peripheral typical network and small dots.

Please cite this article as: Diluvio L, Mazzeo M, Bianchi L, Campanone E. Melanocitosis dèrmica agminada en el territorio del nevus de Ota. Actas Dermosifiliogr. 2018;109:653-655.
The aggregated nevus, \textit{ie} orbital nevus, may appear atypia, epithelial disarrangement.\textsuperscript{5} The extracutaneous assessment to exclude ocular, neurologic and otolaryngologic involvement should be performed routinely. OCT imaging provides optical signs of ocular surface lesions, demonstrating benign findings as hyper-reflective appearance, well circumscribed lesion, regular borders, no invasion of surrounding tissue. Moreover, patients affected by multiple blue nevi, agminated blue nevus or giant blue nevus, particularly caucasians, should be monitored for ocular/orbital involvement and followed closely for signs of rapid growth. Indeed, in blue nevi and uveal melanoma occur somatic mutations in Guanine Nucleotide-binding Protein G(q) (GNAQ), protein encoded by GNAQ gene. These mutations turn GNAQ into a dominant acting oncogene, causing melanocytic neoplasia.\textsuperscript{9,10} We described a case of agminated dermal melanocytosis in the territory of Ota’s nevus, enriching the spectrum of bluish pigmented lesions. Moreover, an integrated team is needed to optimally manage patients with rare neoplasm.

**Conflicts of Interest**

The authors declare that they have no conflicts of interest.

**Acknowledgements**

The authors would like to thank Lucia Pietroleonardo for RCM analysis.

**References**


L. Diluvio,* M. Mazzeo, L. Bianchi, E. Campione
Unidad de Dermatología, Universidad de Roma Tor Vergata, Roma, Italia
*Corresponding author.
E-mail address: lauradiluvio@yahoo.it (L. Diluvio).
1578-2190/© 2017 Elsevier España, S.L.U. and AEDV. Published by Elsevier España, S.L.U. All rights reserved.

Enfermedad de Rosai-Dorfman cutánea: una nueva presentación clínica

Cutaneous Rosai-Dorfman Disease: A Novel Clinical Presentation

To the Editor:

A 21-year-old woman presented with a 3-month history of a diffuse asymptomatic eruption on her face and trunk. She denied fever or any other constitutional symptoms and her review of symptoms was non-contributory. She denied any pertinent past medical or family history. Her only medication included norgestimate/ethinyl estradiol which she had been taking for several years. Physical examination revealed diffuse scattered non-follicular based flesh-colored papules and small nodules. Many lesions demonstrated a central indentation resembling molluscum contagiosum (see Figures 1 & 2). She did not have any cervical, axillary, or inguinal lymphadenopathy. Mucous membrane examination was unremarkable and lacrimal glands did not appear enlarged. Routine histologic examination of her right neck lesion revealed a dense nodular dermal mononuclear cell rich infiltrate showing a significant number of plasma cells and numerous scattered S100 positive multinucleated histocytes with marked emperipolesis and inconspicuous eosinophils (see Figure 3). Complete blood count with differential, erythrocyte sedimentation rate, lactate dehydrogenase, liver transaminases, alkaline phosphatase, bilirubin, and creatinine were negative or within normal limits. Chest and abdominal magnetic resonance imaging were normal. She was treated with a 60 mg oral prednisone taper over six months which resulted in complete resolution of her skin lesions. Follow up at 10 months from onset of disease demonstrated complete remission.

Rosai-Dorfman Disease (RDD) is a relatively rare histiocytic proliferation disorder that was first described in 1969.2 Though RDD classically presents with bilateral massive lymphadenopathy and systemic symptoms, it typically has a benign clinical course and favorable prognosis. In RDD, 43% of patients have involvement of other extranodal sites with skin being the most common site.2 Though skin comprises 10% of extranodal sites involved, approximately 3% are solely cutaneous Rosai-Dorfman Disease (CRDD) without any nodal or other extranodal sites.3 While incidence is reportedly low, CRDD is more prevalent in middle-aged White and Asian women.4 The etiology of RDD is unclear although immunologic,5 viral, and genetic causes including SLC29A3 mutations have been hypothesized.5

The classic presentation of CRDD is a relatively asymptomatic self-involuting nodulo-plaque with surrounding satellite papules.6 However, an evolving wide spectrum of clinical morphologic presentations have been reported. The most common site involved is the face, followed by thigh, and trunk. Recurrence has been reported to occur within 1


Figure 1 Diffuse scattered non-follicular based flesh-colored papules and small nodules, some with central indentations, were seen on the face and the trunk.