Corps Ronds and Corps Grains-Warty Dyskeratoma

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Abbreviations: UV: Ultraviolet; PCR: Polymerase Chain Reaction; DNA: Deoxy-Ribo-Nucleic Acid; RNA: Ribo-Nucleic Acid; MWDS: Multiple Warty Dyskeratoma of the Scalp; HKN: Hair Follicle Specific Keratins

Preface

Graham and Helwig addressed warty dyskeratoma as “isolated Darier’s disease” or “follicular dyskeratoma” in 1954. The nomenclature of warty dyskeratoma was adopted by Szymanski in 1957. “Isolated dykeratosis follicularis” or warty dyskeratoma cognizes an infrequent, cutaneous proliferative lesion commonly situated on the head or neck. Warty dyskeratoma is described as an exceptional, benign skin disease which demonstrates a characteristic solitary papule or nodule with an impacted centroidal crater or comedo [1,2].

Disease Characteristics

Warty dyskeratoma is exemplified by a benign, papulo-nodular lesion which typically demonstrates an endophytic pattern of proliferating squamous epithelial cells, particularly in conjunction with folliculo-sebaceous unit. Preponderant acantholytic foci are additionally enunciated. Frequently, a singular, warty, dyskeratotic papule or nodule with occasional centric crater is cogitated.

Warty dyskeratoma predominately appears in the middle aged to elderly population. The disorder is of obscure aetiology. Ultraviolet (UV) light, autoimmune attributes, exposure to chemical carcinogens and tobacco intake can be implicated in disease pathogenesis.

Viral (human papilloma virus) infection as discerned with a polymerase chain reaction (PCR) assay is absent. However, detection of viral deoxy-ribo-nucleic acid (DNA) and ribo-nucleic acid (RNA) within the cellular epithelium layering villous projections can be cogitated as an incriminating factor in the genesis of warty dyskeratoma.

Chronic actinic deterioration is devoid of implication in the genesis of warty dyskeratoma as lesions can appear in regions unexposed to sunlight [2,3].

An acquired genetic mutation of ATP2A2 gene is discerned along with an absence of SERCA2 gene. Lesions of warty dyskeratoma are engendered from hair follicles as is indicated by preponderance of warty dyskeratoma on hair bearing areas such as scalp. Head and neck are a frequent site of disease incrimination. Lesions do not recede spontaneously.

Warty dyskeratoma can appear on mucosal surfaces at sites devoid of hair follicles. However, inter-relation amidst mucosal and cutaneous lesions remains obscure.

Histology of warty dyskeratoma is identical to keratotic papules of Darier’s disease and warty dyskeratoma can be contemplated as a variant of Darier’s disease.

Individuals with multiple warty dyskeratoma usually lack a family history of adjunct conditions such as Darier’s disease or acantholytic dermatosis as cogitated in Grover’s disease. Multiple warty dyskeratoma can elucidate several or an enhancing quantification of asymptomatic papules on the scalp or forehead. Innumerable, flesh coloured or greyish, hyperkeratotic papules or nodules with a distinctive, umbilicated core can be delineated [3,4].

Multiple warty dyskeratoma of the scalp depicts a female preponderance (70%), a Male: Female ratio of 1: 1.8, a mean age of appearance at 56 years with a disease extent varying from 29 years to 84 years. Lesions can enumerate from 3 to 240.

Multiple warty dyskeratoma of the scalp can display associated morbidities such as cardiac dysarrythmia, hypertension, renal failure and chronic liver disease. Multiple warty dyskeratoma of the scalp is a rare variant and requires a distinction from acquired papulo-nodular lesions in adults.

Multiple warty dyskeratoma of the scalp necessitates a cogent histology, although dermatoscopy can be efficacious in diagnosis [3,5].
Clinical Elucidation

Majority of the lesions are solitary in warty dyskeratoma. Characteristically flesh coloured or brown, verrucose papules, nodules and cysts are enunciated. Exceptionally, multiple. lesions can appear on the face, scalp or dorsum. Adjunctive sites are implicated such as the cheeks, hands and chest.

Frequently manifested is a distinctive, discrete, cup shaped papule or nodule situated on the head and neck or trunk and extremities. Warty dyskeratoma can appear on mons pubis, sub-ungual region, oral and genital mucosa [4,5]. Lesions vary from 1 millimetre to 12 millimetre in magnitude. The disorder is essentially asymptomatic; however, pruritus can occur accompanied by a relapsing, offensive, cheesy effluvium from the lesions. Centric haemorrhage can emerge in traumatic lesions. A centric zone akin to an umbilicus or a pore with a centroidal keratin plug can be cogitated. Multitudinous lesions of warty dyskeratoma are infrequent. Rare, multiple lesions appearing on the scalp are exemplified as “multiple warty dyskeratoma of the scalp” (MWDS). The variant was initially discerned in 1993. Multiple warty dyskeratoma of the scalp depicts an equivocal gender distribution with a Male to Female ratio of 1:1. As denominated, lesions are discerned on the scalp, face, neck, hand or torso. Scalp is a frequent site (75%) followed by face (50%). Asymptomatic scalp lesions are gradually progressive and demonstrate brownish papulo-nodular outgrowths which are minimally pruritic, measure 0.2 centimetres to 2.0 centimetres, evolve gradually and are discerned within months or years. Nodules can be erythematous with a yellowish, hyperkeratotic occlusion. Brownish or black verrucous lesions can be discretely disseminated or clustered [5,6].

Histological Elucidation

Warty dyskeratoma is represented by cup shaped invaginations of the epidermis displaying epithelial hyperplasia and is accompanied by supra-basal acantholysis and dyskeratosis. The morphological characteristics are akin to Darier’s disease. Typical lesions of warty dyskeratoma depict a well-defined, endophytic pattern of evolution with the appearance of corps ronds and corps grains. Additionally, a superficial covering of parakeratotic tissue is delineated. Dyskeratosis is predominantly and characteristically acantholytic. Aberrant linkage and keratinisation of squamous epithelial cells are cogitated. Inverted epidermal invaginations are congested with keratinous debris. Prominent clefts upon supra-basal epithelium are situated inferiorly on epidermal invaginations and configure ascending villous projections. Keratinous deposits are abundant and articulate keratin plugs with centric cellular proliferation [6,7].

Innumerable villous extensions upon the inferior segment of cup shaped depression are coated with a singular layer of basa-loid epithelial cells. Invaginations can appear within an intensely distended hair follicle. A singular lesion can implicate two to four adjoining hair follicles. Admixture of histological features can be enunciated. Dyskeratotic keratinocytes appear within the stratum spinosum and stratum granulosum with the elucidation of corps ronds and corps grains, respectively. An accompanying thickened granular layer is demonstrated. Corps ronds and corps grains can project through the papillary dermis. Centric crater is distended with necrotic, cornified tissue. Sequential to the frequently elucidated cup shaped invaginations, cystic and nodular configurations are additionally cogitated.

Cystic category depicts enlarged; well demarcated cystic articulations confined to the dermis. Cystic arrangements are layered with epithelial cells and impacted with abundant keratinous substance. Nodular variant demonstrates miniature, well defined, circumscribed, solid aggregates of epithelial cells within the dermis. Epithelial accumulates are in continuation with superficial epidermis which enunciate epidermal hyperplasia [7,8]. Minimal to moderate inflammatory infiltrate comprising of lymphocytes and histiocytes can accrue in the superficial dermis underlining the tumefaction. Focal emergence of acantholytic dyskeratosis cannot be ascertained as a categorical morphological parameter of warty dyskeratoma.

Focal acantholytic dyskeratosis is additionally discerned in Darier’s disease, transient acantholytic dermatosis, acantholytic squamous cell carcinoma, actinic keratoses and basal cell carcinoma. Multiple warty dyskeratoma of the scalp on histology depict a distinct, predominantly endophytic epidermal proliferation constituting a centroidal crater with impacted keratinous substance composed of dyskeratotic keratinocytes which additionally configure the classic corps ronds and grains. Inferior aspect of the principally epidermal lesion exhibits acanthosis, prominent papillomatosis and accumulation of dyskeratotic keratinocytes which appear submerged in acantholytic, supra-basal epithelial perforations. Epidermal extensions simulating a “villous” arise within epidermal clefts. Typical histology, abutting hair follicles and representative immune histochemical reactivity to hair follicle specific keratins (HKN) 5/17, particularly HKN-6 and HKN-7 is diagnostic. Thus, an alternative terminology of “follicular dyskeratosis” can be cogitated for multiple warty dyskeratoma of the scalp [8,9].

Dermoscopic Features

Dermoscopic manifestations of warty dyskeratoma include whitish, homogenous zones demonstrating yellow cods interspersed with mature, distinct hair follicles. Additionally, pale to gray zones with a centric depression can be elucidated on dermascopy [3,4].

Differential Diagnosis

Differentiation is necessitated from clinical conditions such as verruca, seborrheic keratoses, sebaceous cyst, sebaceous hyperplasia and hypertrophic actinic keratoses. Histological demarcation is a pre-requisite from conditions demonstrating acantholytic dyskeratosis on morphology such as acantholytic or dyskeratotic acanthoma, Darier’s disease, Grover’s disease, Hailey- Hailey disease, familial dyskeratotic comedones, acantholytic actinic keratoses and acantholytic squamous cell carcinoma.
Appearance of definitive clinical aspects and emergence of villous epidermal hyperplasia at inferior segment of warty dyskeratoma assists in segregation from identical morphologies [9-11]. Warty dyskeratoma can be differentiated from Darier's disease and transient acantholytic dermatosis on the basis of specific clinical attributes. As the lesions are devoid of cellular atypia and display particular morphological features, conditions such as squamous cell carcinoma, actinic keratosis or basal cell carcinoma can be excluded. Differentiation on histological grounds is necessitated from conditions such as i) Benign familial pemphigus or Hailey-Hailey disease which displays excessive acantholysis within enlarged zones of epidermis. ii) Darier’s disease which requires clinic-pathological concordance for appropriate diagnosis. iii) Pemphigus vegetans which demonstrates enhanced foci of acantholysis. Eosinophilic aggregates are cogitated within the epidermis. Surgical specimens submitted for immune fluorescence can be utilized for discerning pemphigus vegetans [3,4] [Figures 1-12].

Therapeutic Regimens
Optimal treatment comprises of surgical extermination of the lesions, although it is not a pre-requisite with benign histology. Surgical eradication is a cogent form of therapy. Partial or incompletely elucidated lesions can be suitably surgically eliminated.
Apart from surgical excision, curettage, electrodessication and irradiation with X rays can be employed to remove the lesions. However, lesions can reoccur with the utilization of aforesaid treatment modalities.

Administration of tazarotenic acid gel is beneficial. Systemic administration of isotretinoin in combination with topical tretinoin application can achieve significant alleviation. Reoccurrence of the lesions is exceptional. Malignant transformation does not ensue [9,11].

References


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