Down Syndrome and Delayed Occurrence of Acne

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Down syndrome (DS) is associated with an increased incidence of skin diseases which may cause further psychosocial disability in affected patients [1, 2]. In this setting, an increased incidence of acne has been associated with DS in some previous publications [1]. Some data may support such an association. The use of anticonvulsive drugs and an altered zinc metabolism in patients with DS [3] might promote the development of acne. Overweight is also common in DS patients [4]. However, although some recent studies suggest that a high-glycemic-load diet may facilitate acne [5–8], a link between a high BMI and the development of acne is far from being proved [8].

In this issue of Dermatology, Bagatin et al. [9] found that DS patients have a low prevalence of acne, suggesting that trisomy 21 may have an unexpected protective effect against acne. Interestingly, the prevalence of acne was higher in DS patients older than 19 years than in teenagers, suggesting a delayed occurrence of acne. The authors hypothesize that, considering the possible link between stress and acne, delayed neuropsychomotor development in DS patients could lead to less psychological stress. However, despite their well-known cheerful and friendly demeanor creating a personality stereotype, approximately 20–40% of children with DS display recognized behavioral problems [10].

Several endocrine abnormalities are likely to contribute to a low prevalence of acne in DS patients. Androgens, such as dihydrotestosterone and testosterone, the adrenal precursor dehydroepiandrosterone sulfate, estrogens such as estradiol, and other hormones, including growth hormone and insulin-like growth factors (which are maximally secreted during adolescence), may each be important in the development of acne [11]. DS patients are known to suffer from growth hormone deficiency and inadequate insulin-like growth factor response, resulting in remarkable skeletal maturation delay and short stature [12]. DS patients also have a high incidence of abnormalities in their sexual development, and delayed puberty is reported in both sexes. Reported abnormalities in girls include hypogonadism with a delay in either menarche or adrenarche. In boys, described defects vary from ambiguous genitalia, cryptorchidism, micropenis, small testes and low sperm count to scant development of axillary hair and beard [12]. Mean levels of serum follicle-stimulating hormone and luteinizing hormone are usually markedly elevated, but mean testosterone levels are normal, which suggests a diagnosis of partial primary gonadal dysfunction [13]. Taken together, these data suggest that, despite premature aging, the onset of acne is delayed in DS patients, and support the relationship between acne and pubertal age rather than chronological age.
References