The dysplastic naevus

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Since the dysplastic naevus (DN) syndrome was first described, it is marker lesion, the DN, has been subject of vehement debate. Major disagreements concerning its macroscopic and microscopic characteristics, its prevalence, its relation to melanoma, its treatment, and indeed its very existence as a separate entity, have continued for decades. As a result, some have dismissed the entire matter as unsolvable. Some now prefer alternative or related terms such as atypical mole or an eponymic designation, or have included severely dysplastic naevi in a category of melanocytic intrapidermal neoplasia (MIN). Others have continued to maintain that the dysplastic naevus is a distinct entity, which can and should be distinguished from other naevus types and from melanoma.

Here, an attempt is made to analyse some of the causes of this controversy, with special emphasis on the relation between the predictive value of diagnostic criteria of entities entering the differential diagnosis and the relative prevalence of these entities. Arguments in favour of the continued use of DN as a separate entity will be provided, and practical guidelines for its diagnosis will be summarised.

Usefulness of the terms dysplasia and DN

The term dysplastic naevus has been repeatedly criticised. In pathology, the term dysplasia is used in very different ways—for example, dysplastic goiter, dysplasia of the uterine cervix, bronchopulmonary dysplasia. Therefore, is was argued, the term would be too ambiguous to be used at all. However, it should be borne in mind that in all language, including scientific language, words have different meanings in different contexts. The context defines its meaning in the individual instance. Indeed, the word mole may refer to an animal or a cutaneous pigmented lesion. The term dysplasia in the context of naevi indicates a combination of cellular and architectural irregularities, described in detail below, and associated to varying degrees (depending primarily on clinical context) with increased melanoma risk.

Nonetheless, some dermatologists and pathologists now favour alternative designations such as atypical mole. However, the words atypical and mole are at least as varied in their meaning as is the word dysplastic. The term melanocytic intraepidermal neoplasia (MIN), which lumps together severely dysplastic naevi and in situ melanoma, has the advantage of abolishing the problem ridden histological distinction between DN and in situ melanoma, a distinction that has no apparent clinical relevance, but has other disadvantages. Very severely dysplastic DN are compound rather than intraepidermal, and the term neoplasia would also be applicable to other naevus types. In my opinion, the term DN remains the best choice.

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should not be made when the lesion is smaller than 5 mm in diameter clinically or 4 mm across on the slide. This is because a large number of small naevi, at an early stage of their development when still only a few millimetres in diameter, exhibit some or many of the histological features of DN, especially irregular nesting with occasional rete ridge bridging, as well as lamellar subepidermal sclerosis. It is the combination of larger size and the histology that characterises DN.

**ARCHITECTURAL FEATURES**

Most DN are compound, possessing a junctional and an intradermal component. A small minority of DN are entirely junctional. No intradermal DN is recognised. Compound dysplastic naevi often have a slightly raised centre and a flat periphery, which consists of junctional lateral spread of the proliferation, associated with irregular rete ridge elongation and subepidermal sclerosis (fig 1), often consisting of stacks of slightly refractile collagen fibres (lamellar fibrosis; fig 2).

The junctional component generally consists of nests as well as solitary cells arranged in a lentiginous pattern. The nests are irregularly sized and shaped, and often have a horizontal orientation bridging adjacent epidermal rete ridges. The lentiginous proliferations of melanocytes are similarly irregularly distributed (fig 1). The junctional component is highly cellular, in areas replacing the large majority of basal epidermal keratinocytes. Both nests and lentiginous proliferations usually extend somewhat along the epithelium of cutaneous adnexae. Mitotic figures are rare. Ascent of solitary melanocytes so that these cells reach the granular layer is also absent or rare in DN.

If present, the ascending cells are small, with compact, dark nuclei, and little cytoplasm.

The dermal component of dysplastic compound naevi is located at the centre, so that the lesion is symmetrical. Generally, well defined nests and strands of melanocytes predominate in the subepidermal part, which exhibits distinct sclerotic changes. Deeper parts of the naevus consist of ill defined aggregates or, less commonly, more compact small groups of cells. Larger, compact, spherical intradermal nodules are absent—their presence should raise a strong suspicion of (vertical growth phase) melanoma. Melanocytes at the base of the lesion are usually small but may be pigmented. Generally, the intradermal component of DN is not very extensive.

The dysplastic naevus exhibits a characteristic host response, consisting of irregular rete ridge elongation, together with subepidermal sclerosis, which takes the shape of dense, refractile stacks of collagen fibres around rete ridges and along capillaries of dermal papillae. Proliferation of dermal capillaries and a perivascular lymphohistiocytic inflammatory infiltrate are invariably present.

Importantly, the architectural and cellular features of different parts of the naevus are similar—at any level of the lesion, the cell type is similar throughout. In this, the DN resembles most other naevus types, and is distinct

**Morphology of the DN**

The macroscopical features of the DN are best evaluated before removal. DN are roughly symmetrical, flat or slightly raised pigmented lesions, or they may have a slightly papular centre surrounded by a flat rim. Most are over 6 mm in diameter. DN are not verrucous or pendulous. The peripheral border is often blurred and slightly irregular, and the lesion is surrounded by a reddish hue, caused by reactive hyperaemia; this feature is no longer evaluable after excision. The pigmentation may be slightly irregular in intensity and colour. In DN syndrome patients, DN are often numerous and occur in sites where naevi are generally absent or scarce, such as the buttocks, breasts, genital skin, and dorsa of feet. Occasionally, a DN syndrome patient may have fewer than 10 clinically evident DN.

The histological diagnosis of DN is based on a combination of cytological and architectural features, including a characteristic host response. It is advised that the diagnosis of DN

**Figure 1** Dysplastic compound naevus. Irregular distribution melanocytes arranged in nests and lentiginous patterns along the dermoepidermal junction. Irregular rete ridges, in part surrounded by lamellar sclerosis. Slightly irregular distribution of intradermal melanocytic nests and smaller cell groups.

**Figure 2** Dysplastic compound naevus. Irregular rete ridges, surrounded by pronounced lamellar sclerosis.
The dysplastic naevus

from most melanomas, which exhibit more variation in architecture and cellular features.

CYTOLOGICAL FEATURES
The melanocytes of DN exhibit nuclear atypia, especially nuclear pleomorphism and anisochromatism, most obvious at the dermoeidermal junction and in the subepidermal region (fig 3). Such nuclear atypia is a prerequisite for a positive diagnosis of DN. It is the variability in size, shape, and staining intensity of the nuclei, rather than their absolute size, that is most characteristic of DN. Nuclei of melanocytes in Spitz naevi, pigmented spindle cell naevi, blue naevi, and deep penetrating naevi may be large, but in these latter naevus types, the variability of nuclear features is less striking than in a DN. In contrast to most melanomas, the nuclear pleomorphism is similar in different areas of the lesion. In melanoma, there is often a difference in nuclear features between different nests or aggregates.

In my experience, the presence of dusty pigment in junctional melanocytes is of little practical use in the differential diagnosis, as it may also be present in melanoma as well as in some common acquired naevi, Spitz naevi, deep penetrating naevi, and balloon cell naevi. Intradermal mitotic figures are absent or exceedingly rare. Their presence should raise a suspicion of melanoma.

None of the histological features of DN that have been put forward is diagnostic by itself, it is a combination of features, rather than any single feature in isolation, that sets apart these naevi. If used in this way, there is a fair correlation between the histological and clinical diagnosis of DN.

As each of the features relevant to the diagnosis of DN can be present to various degrees, and in various combinations, classification problems of borderline cases are unavoidable. This is especially problematic with respect to the differential diagnosis between common acquired naevus and DN.

Obviously, the combination of distinct cyto- logical and architectural atypical features in a naevus of an affected familial DN syndrome patient does not provide major diagnostic difficulties. However, such cases are rare in comparison to the very large number of naevi with only some of these features, occurring in otherwise unremarkable individuals. It is clear that a restrictive diagnostic approach to DN will exclude the large majority of these latter lesions.

The grading of dysplasia in DN is advocated by some workers; however, since it results in significant intraobserver and interobserver variability, I have discontinued grading DN.

Relative prevalence of entities v predictive value of diagnostic criteria
An important consideration, often overlooked in discussions on the practical value of diagnostic criteria, is the profound impact of the relative prevalence of two entities requiring differentiation, on the predictive value of a given criterion used to differentiate the two. In other words, a feature, or a combination of features, needs to have a very high specificity if it is to be of use in the positive identification of a rare entity.

This is probably of central relevance to the controversy on dysplastic naeves. Naevi submitted for histology constitute a very small proportion of the naevi in the general population. The reason for removal is usually either cosmetic or a suspicion of melanoma. Slides referred for expert opinion reflect a substantial bias, the numbers of problem cases, including DN, is much higher in such series. Significantly, published series used for the study of interobserver variability in the diagnosis of DN have included relatively large numbers of DN, which contrasts with the material received in general diagnostic pathology laboratories where non-dysplastic naevi greatly outnumber DN. In these latter diagnostic pathology laboratories, a problem of insufficient specificity of diagnostic criteria for DN is encountered; this problem is masked to a considerable extent in studies based on selected material in which DN are greatly overrepresented. To avoid this problem, as far as possible a conservative approach in the diagnosis of DN is advised: only those naevi 6 mm or larger in diameter and meeting the combination of the cytological and architectural abnormalities summarised above should be designated DN.

Dysplastic naevus v early melanoma
The distinction between DN and melanoma is of greater importance clinically than between DN and common acquired naevi. A number of negative findings is of central importance in the distinction of DN from small melanomas. In DN, there is no pagetoid spread of atypical melanocytes at all levels of the epidermis, across most or all of the lesion. Moreover, intradermal mitoses are absent, unless a brisk inflammatory infiltrate permeates the dermal part of the lesion. Atypical mitotic figures are invariably absent. Distinct differences in pres-
ence, degree or aspect of inflammation and fibrosis between different areas of the lesion are lacking in DN.

The malignant potential of many thin, presumably early, melanomas as they are diagnosed today is far from clear. The question arises: are these so-called early melanomas really melanomas—that is, fully malignant tumours that were discovered in time, or do they only resemble melanoma, clinically and histologically, but need further steps before the potential to invade and metastasise is acquired? Follow up data of the large material of Elder et al.27 and Clark et al.28 indicate that radial growth phase melanomas never metastasise. Although independent confirmation of their conclusions is necessary before the acceptance of radial and vertical growth phase melanoma as diagnostic concepts, their findings raise an interesting point: if there is a phase in which a melanoma never metastasizes, it would seem to be appropriate to conclude that at this stage the lesion may be a precursor lesion of malignancy, but has not yet acquired a fully malignant phenotype, and therefore is not really an early melanoma. Further studies on thin metastasising melanomas are essential to identify small melanomas with a greater degree of precision, and to attain a greater degree of confidence in excluding the diagnosis of melanoma in thin worrisome lesions. The outcome of such ongoing work will no doubt contribute significantly to the distinction between DN and melanoma.

The practical role of the pathologist in the diagnosis of dysplastic naevi

Early reports on DN seemed to indicate that the histopathologist would be able to identify with considerable reliability a naevus type that, regardless of clinical context, would constitute a marker of clinically significant melanoma risk. Clearly, this is not the case. It is the clinical evaluation (numbers, distribution and macroscopic appearance of the patient’s naevi, and family history) that is the best basis for the diagnosis and clinical management of the DN syndrome. It should, however, be borne in mind that the clinical reproducibility of DN, as shown by recent interobserver variability studies based on photographs of single naevis, is limited.29 30 The primary task of the histopathologist, with respect to DN, is to rule out melanoma. A patient with the clinical phenotype of the DN syndrome should be carefully monitored and excessive exposure to sunlight should be avoided. Prophylactic removal of all DN is a futile exercise, DN continue to arise, change in appearance, and disappear throughout adult life.31 32 A change in size or appearance of a previously stable naevus raises a suspicion of malignant transformation and should lead to excision of the lesion. The practical relevant question in such an instance is not, is the naevis dysplastic? but rather, is the lesion a melanoma?

If the pathologist diagnoses a DN, it is wise to add a note in the report indicating that this finding, in itself, does not imply that the patient has a significant risk for melanoma, but that a proper clinical evaluation of the numbers, distribution, and appearance of naevis across the entire skin is advised. If the clinical findings are within normal limits, the diagnosis apparently has no obvious further consequences.

Conclusions

From the above, a number of conclusions can be drawn:

(1) The clinical diagnosis of familial DN syndrome is based primarily on the clinical assessment of number, distribution, and appearance of naevi on the entire skin, in combination with family history.

(2) The histological investigation of a single naevus does not effectively identify DN syndrome patients. The main objective of the histological investigation of a naevus in the clinical setting of DN syndrome is to rule out melanoma.

(3) Substantial controversy in the histopathological literature concerning the predictive value of criteria or sets of criteria for the diagnosis of DN is probably related to large differences in relative prevalence of entities in series studied or materials received in the diagnostic pathology laboratory.

(4) A restrictive approach in the histological diagnosis of DN is advised, as it results in a better correlation with personal or family history of melanoma.

In recent years, genetic studies of familial DN syndrome kindred have resulted in the identification of at least two loci, on chromosomes 1 and 9,33 associated with the DN syndrome phenotype. Apparently, there is genotypic heterogeneity in this syndrome. Future studies of the genetics of DN syndrome may provide clues at the DNA level, which may hopefully constitute an additional marker in the pathological diagnosis of sporadic DN.

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