Histological features of CIN3 and their value in predicting invasive microinvasive squamous carcinoma

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Abstract

Aims—To determine the histological features in CIN3 associated with or predictive of subsequent microinvasion.

Methods—The histological appearances of CIN3 accompanying 120 cases of microinvasive carcinoma of the uterine cervix were retrospectively studied. Major features were defined as those present in greater than 80% of cases of microinvasive carcinoma (MICA) and less than 10% of control cases of CIN3. One hundred cases of CIN3, 36 showing all, and 64 lacking all, of the major features associated with microinvasion, as defined in the retrospective study, were prospectively studied. Deeper levels were cut to exclude the presence of microinvasion in the original biopsy specimen and negative cases were followed up for a period of up to 18 months in order to assess rates of recurrence or progression.

Results—The major features identified in CIN3 associated with microinvasive carcinoma were extensive involvement of surface epithelium and deep endocervical crypts by expansile CIN3, luminal necrosis, and intraepithelial squamous maturation. Other features more commonly present in MICA associated CIN3 than in controls included frequent mitosis and apoptosis, pericryptal concentric fibroplasia, pericryptal inflammatory infiltrate, pronounced cellular pleomorphism, nuclear changes (distinct nucleoli and chromatin clearing), and the emergence of streams of darkly stained spindle cells orientated at right angles to the basement membrane. In the prospective study 83% of cases illustrating the major MICA associated features revealed evidence of MICA or frank invasion either on serial sections of the original biopsy or on subsequent biopsy. None of the 64 cases of CIN3 that lacked these features showed evidence of invasion on serial sections or on further follow up over 18 months.

Conclusions—The data strongly suggest that cases of CIN3 which have a higher probability of association with or rapid progression to invasive disease can be identified. When these features are present in a biopsy specimen of CIN3, serial sections should be performed to exclude the presence of microinvasion. Closer clinical follow up of these patients may be needed.

Most cases of invasive squamous carcinoma of the cervix are thought to be preceded by the metaplasia-intraepithelial neoplasia (CIN) sequence. This has formed the basis for cervical screening programmes to detect the earlier stages of the disease in the hope of preventing invasive carcinoma and reducing the mortality from the disease. Most surgeons currently treat all cases of CIN with loop diathermy, laser, or cold coagulation therapy and follow them up regularly by cervical smears and colposcopic assessment. In spite of these measures some women will still have recurrence or progression of their disease. The reasons why some cases of CIN and in particular CIN3 recur or progress after treatment are currently uncertain. One of the reasons may be that the original lesion has not been fully excised, or a minute focus of microinvasive carcinoma (MICA) has been left behind or has not been detected in the biopsy specimen either due to misinterpretation or inadequate sampling. Another possibility is the existence of a biologically more aggressive subtype of CIN3. If this assumption is true it would be of paramount importance to identify the subtype. There is some indication, on the basis of cytogenetic studies, that CIN3 is heterogeneous in terms of risk of progression to cancer. On the basis of morphology alone, however, there is currently no recognised way of predicting which cases of CIN3 will progress. It is also controversial whether HPV associated CIN is biologically more sinister, in particular CIN containing HPV 18 rather than 16.

The Department of Pathology at Edinburgh University receives about 4000 colposcopic punch and loop biopsy specimens annually. Of these, 0-4% show MICA and 0-8% show invasive carcinoma. At weekly combined oncology/gynaecology/pathology meetings the management of all cases of invasive and microinvasive squamous carcinoma are discussed and current and previous histological samples reviewed.

On review of previous biopsy specimens from cases showing recurrence or progression and CIN3 adjacent to microinvasion, similar histological features were often noted. The aims of this study were to identify histological...
appearances in cases of CIN3 showing a significant association with microinvasion, and to test their utility in predicting subsequent microinvasion in a prospective series of cases of CIN3.

Methods
PART I: RETROSPECTIVE STUDY OF FEATURES ASSOCIATED WITH MICROINVASIVE CARCINOMA
One hundred and twenty cases of microinvasive squamous carcinoma of the uterine cervix and 80 control cases of CIN3 without microinvasion were retrieved from the departmental surgical files made between 1988–1992. All were from loop excisions of cervix which had been fixed in Bouin’s solution and serially cut into six to 20 parallel blocks of about 3 mm in thickness. These were routinely processed and paraffin wax sections were stained with haematoxylin and eosin.

Microinvasive carcinoma
The criteria used for the diagnosis of MICA were those proposed by the Society of Gynaecological Oncologists (SGO) in 1974 and modified in 1985: “MICA is defined as a lesion that invades the stroma to a depth of 3.0 mm or less, and in which there is no evidence of lymphatic space invasion”.

SGO classification rather than FIGO was used because in a recent literature review, it was shown that the difference in the incidence of lymph node metastasis for tumours 0.1 to 3.0 mm and for tumours 3.1 to 5.0 mm in depth is significant (0.7% vs 4.3%) and therefore a small number of women with FIGO stage 1a 2 (up to 5 mm in depth and 7mm in width) will have tumour extension beyond the uterus.

Foci of microinvasion were identified by squamoid differentiation in groups of cells separated from or protruding through the basement membrane of areas of CIN3 (fig 1). The depth of stromal invasion was measured using an ocular micrometer; the measurement was made between the deepest part of the invasive focus and the basement membrane of the overlying epithelium.

Overlying/adjacent CIN3
The following histological features were evaluated:
1. The extent of the lesion.
2. The proportion of crypts or surface affected by the lesion. Greater than about 25% of the glands or surface epithelium included in the specimen was considered to be extensive disease.
3. The complexity of the architecture of the lesion.
4. Expansion of the affected crypts or surface epithelium.
5. Secondary features.
6. The presence or absence of central (comedo-like) necrosis or inflammation.
7. The cell morphology.
8. The presence or absence of easily identifiable apoptosis or single cell dyskeratosis, the degree of pleomorphism and mitotic activity, and the nuclear pattern (presence of chromatin clearing or acquisition of nucleoli).
9. Appearance of adjacent stroma (fibrosis, inflammation).

The significance of the association of each of the above features with MICA evaluated using the χ² test.
Table 1  Incidence of various histological features found in association with microinvasive squamous carcinoma of cervix

<table>
<thead>
<tr>
<th>Histological features</th>
<th>Control cases (total 80 cases)</th>
<th>CIN3 with MICA (total 120 cases)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surface involvement only</td>
<td>34 (43%)</td>
<td>9 (7%)</td>
</tr>
<tr>
<td>Involvement of deepest endocervical crypts present in specimen</td>
<td>1 (1%)</td>
<td>111 (93%)</td>
</tr>
<tr>
<td>Irregular expansion of involved surface/crypts</td>
<td>5 (6%)</td>
<td>106 (96%)</td>
</tr>
<tr>
<td>Necrotic debris</td>
<td>4 (5%)</td>
<td>99 (82%)</td>
</tr>
<tr>
<td>Intralesional maturation</td>
<td>4 (5%)</td>
<td>105 (87.5%)</td>
</tr>
<tr>
<td>Abnormal mitoses</td>
<td>20 (25%)</td>
<td>53 (44%)</td>
</tr>
<tr>
<td>Apoptosis</td>
<td>13 (16%)</td>
<td>101 (84%)</td>
</tr>
<tr>
<td>Dyskeratotic cells</td>
<td>11 (14%)</td>
<td>67 (55%)</td>
</tr>
<tr>
<td>Nuclear clearing with prominent nucleoli</td>
<td>6 (8%)</td>
<td>67 (55%)</td>
</tr>
<tr>
<td>Pericryptal concentric fibrosis</td>
<td>8 (10%)</td>
<td>75 (63%)</td>
</tr>
<tr>
<td>Pericryptal lymphocytic infiltrate</td>
<td>20 (25%)</td>
<td>92 (77%)</td>
</tr>
<tr>
<td>Intralesional lymphocytic infiltrite</td>
<td>17 (21%)</td>
<td>78 (65%)</td>
</tr>
<tr>
<td>Emergence of small dark spindle cells</td>
<td>12 (15%)</td>
<td>34 (28%)</td>
</tr>
</tbody>
</table>

All of the above criteria were significantly associated with the presence of MICA (p < 0.001) except the emergence of darkly staining spindle cells (p = 0.05). In the case of restriction of disease to the surface epithelium, this association was significantly negative.

Results

PART I: RETROSPECTIVE STUDY

Distribution of the lesion

Extensive involvement (more than 25% of epithelium) of both surface epithelium and endocervical crypts, including the deepest crypts present in the specimen was seen in 93% of cases of MICA and 1% of controls.

Architecture of the involved epithelium

In 96% of cases of MICA the involved crypts were distinctly expanded by the presence of CIN3, often with irregular bulging of their borders (fig 2). Similarly, the affected surface epithelium in these cases was irregularly expanded but to a lesser degree than the crypts. These features were present in 6% of controls.

Necrosis

Comedo-like central necrosis consisting of desquamated dyskeratotic cells admixed with polymorphs was seen within the lumen of the affected crypts with analogous necrosis of the surface epithelium in 82% of cases of MICA and 5% of controls (fig 3).

Intralesional maturation

Focal intraepithelial squamous maturation in the form of whirling, abortive squamous eddies, and cytoplasmic keratinisation was seen in 86% of cases of MICA and 5% of controls (figs 3 and 4).

Cytological patterns

The constituent cells exhibited mild, moderate, or severe cellular pleomorphism (fig 5). Some of the nuclei showed coarsening of the chromatin, focal clearing, and prominent nucleoli (55% of cases of MICA, 8% of controls). Isolated single dyskeratotic cells were seen in 55% of cases of MICA and 14% of controls and were often associated with numerous apoptotic cells (84% of MICA and 16% of controls). Varying numbers of normal mitotic figures were present in all cases, and abnormal forms were seen in 44% of cases of MICA and 25% of controls. Finally there were streams of darkly stained, spindle-shaped cells compressed inbetween the remaining epithelial cells and these seemed to be orientated vertically from the basement membrane towards the luminal surface (fig 6). These were seen more often in the invasive foci themselves but also in the overlying abnormal epithelium in 28% of cases of MICA. These cells were also identifiable in 15% of controls. Scattered intraepithelial lymphocytes were found in 65% of cases of MICA and 21% of controls.

Adjacent stroma

A concentric lamellar fibrosis was seen around the crypts in 63% of cases of MICA and 10% of controls (fig 7), and a moderate or dense lymphocytic infiltrate was shown in 77% of cases of MICA and 25% of controls.

PART II: PROSPECTIVE STUDY (table 2)

None of the 64 cases of CIN3 that lacked the
Figure 3 CIN3 involving and expanding endocervical crypt (A). Note intralesional squamous maturation and the central dyskeratotic necrosis (B) (haematoxylin and eosin).

Table 2 Comparison of prospectively studied cases showing or lacking major MICA associated features

<table>
<thead>
<tr>
<th>Incidence of features</th>
<th>CIN3 with all major (n = 36)</th>
<th>CIN3 with no major (n = 64)</th>
</tr>
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<tbody>
<tr>
<td>Coexistent MICA</td>
<td>10 (28%)</td>
<td>0</td>
</tr>
<tr>
<td>Recurrence as CIN3</td>
<td>1 (3%)</td>
<td>0</td>
</tr>
<tr>
<td>Recurrence as MICA</td>
<td>10 (28%)</td>
<td>0</td>
</tr>
<tr>
<td>Recurrence as invasive carcinoma</td>
<td>9 (25%)</td>
<td>0</td>
</tr>
<tr>
<td>Incomplete excision in original biopsy specimen</td>
<td>29 (80%)</td>
<td>33 (52%)</td>
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</table>

Figure 4 CIN3 lesion. Note intralesional squamous differentiation and early squamous eddies (arrow). A tongue of impending invasion is also seen (haematoxylin and eosin).

Figure 5 CIN3 involving endocervical crypt. Note cellular pleomorphism, apoptosis, abnormal mitoses and central necrosis (haematoxylin and eosin).

Discussion

Effective cytology screening programmes have meant that carcinoma of the uterine cervix is being detected at an earlier stage in its evolution and fewer women are now presenting with advanced cancer. In achieving this success, the screening programme has led to a huge increase in surgical interventions, many of which may arguably represent overtreat-
Figure 6 CIN3 showing pericryptal condensation of cells and the emergence of small dark spindle cells traversing from the basal layers towards the crypt centre (haematoxylin and eosin).

Figure 7 CIN3 involving crypts. Note lamellar fibrosis (haematoxylin and eosin).

ment; yet cases of invasive squamous carcinoma still occur in women who have been regularly screened. Until now it has not been considered possible to distinguish cases of CIN which have a higher risk of rapid progression to invasive disease. If such a distinction could reliably be made on histological grounds, clinical management protocols could be adjusted accordingly, with benefits to both patients with "high risk" or "low risk" disease.

In this study we identified several histological criteria which are present in most (over 80%) of cases of CIN3 accompanying MICA and in a minority of control cases (under 10%), which had predicted aggressive behaviour in prospectively collected cases. The major predictive criteria (table 1) were extensive involvement of both surface and deep endocervical crypts, expansion of the crypts by CIN3, comedo-like central necrosis, and intraluminal squamous maturation. Increasing involvement of the endocervical crypts seems to occur with progression of CIN. Anderson and Hartley found that 88.6% of CIN3 had some crypt involvement; Abdul-Karim et al found that the higher the grade of the CIN the greater the extent of the surface and crypt involvement, and Demopoulos et al have shown that deep endocervical gland involvement by CIN3 is a highly significant predictor of residual or recurrent disease. Another important feature highlighted by the present study was distension or expansion of the involved endocervical crypts or surface epithelium. The frequent finding of comedo-like central necrosis may represent a further progression of this process.

Changes in the cytological appearance of CIN3 adjacent to microinvasive carcinoma have been reported before. These include: islands of well differentiated squamous cells present at all levels of the epithelium; disorganised cellular polarity; cellular pleomorphism; presence of nucleoli in some cells; frequent pyknosis; and individual cell keratinisation. All of these features were confirmed in the present study, but squamous maturation (either in the form of keratinisation or squamous eddies) was found to be the most frequent change. The other cytological changes described above (termed minor criteria in table 1) were frequently present in cases with microinvasion, but were not found to have predictive value in the prospective study.

Whether these changes precede or develop simultaneously with MICA cannot be ascertained with certainty by this study. If they precede MICA this is of great importance, as their presence in CIN3 may indicate incipient invasion. In the second part of this study follow up biopsy specimens from the cases of CIN3 exhibiting all of the major criteria showed MICA in 48% and frankly invasive carcinoma in 23%. One third of the remaining 29% illustrated foci of MICA on serial sections of the original biopsy specimens and the remainder showed no evidence of invasion, either on deeper levels or on subsequent biopsy specimens. This lack of progression may represent adequate treatment, rather than lack of aggressive potential.

None of the cases of CIN3 selected as a control group (lacking any of the major criteria) progressed during the period of the study. Interestingly, retrospective examination of the surgical files over the period of the study showed that none of the surgically treated cases of CIN3 received by our department but not included in this study had progressed, despite CIN3 extending into the endocervical resection margin in 51% of these cases. This suggests that incompleteness of excision may not be the only factor involved in the high rate of recurrence or progression seen in the cases with the major MICA associated features.

The results of this study suggest that there are certain histological features which when present in CIN3 predict a greater risk of progression to invasion, compared with cases lacking these features. We do not propose readoption of the former division of CIN3 into severe dysplasia and carcinoma in situ on this basis, but we feel that when present, the
possible meaning of these features should be communicated to the clinician. The reason for this is twofold. Firstly, even with thorough sampling, it is possible to miss foci of microinvasion, as has been coincidentally shown in the course of this study. Secondly, the high risk category of CIN3 involves endocervical glands more extensively and is therefore more likely to be incompletely excised during standard loop excision, as has been suggested by the study of Demopoulos et al. Clearly, if residual disease of this type progresses more rapidly or is already associated with MICA at the time of surgery, recurrence as more advanced invasive disease (as happened in six cases in this study) is a serious possibility.

We therefore feel that there is a case for using terminology such as “CIN3 with features suggestive of incipient microinvasion,” or “CIN3 with features associated with microinvasion,” in reporting cases of CIN3 showing the high risk pattern described in this study. This would at least instil in the clinician’s mind a requirement for closer follow up (perhaps with endocervical brush cytology) or further excision, particularly if the completeness of the original excision was in doubt.

In conclusion, this study highlights the potential importance of recognising certain features in CIN3 which suggest incipient microinvasion. The major criteria identified were: extensive, expansile involvement of both surface and crypt epithelium, including the deepest crypts present in the specimen; central dyskeratotic necrosis; and intralesional squamous maturation. Other features frequently present in these cases were dyskeratosis and apoptosis, pericytystal lymphocytic infiltration, and fibrolamellar fibrosis, but these features were not found to have predictive value. Whether the high risk cases represent a biologically distinct subset (for example, with relation to human papillomavirus infection) or simply more advanced disease will require further elucidation. In practice, however, we recommend that cases showing all of the major criteria described above should be examined at multiple levels to exclude microinvasion, and that the possible meaning of these histological features should be communicated to the clinician.

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