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Abstract

Aim—To determine the interobserver variation in scoring presence and grade of vulvar intraepithelial neoplasia (VIN) in haematoxylin/eosin (H/E) slides, MIB 1 slides, and the combined use of H/E and MIB 1 slides.

Methods—10 slides were stained with H/E and MIB 1 with each of the following diagnoses: normal vulvar skin, VIN 1, VIN 2, and VIN 3. Six observers first scored the H/E slides separately from the MIB 1 slides and second the combined H/E and MIB 1 slides.

Results—Unweighted group $k$ for MIB 1 was 0.62 and the weighted group $k$ was 0.91. This was significantly better than the unweighted group $k$ for H/E slides (0.47, $p = 0.023$) as well as the weighted group $k$ for H/E slides (0.82, $p = 0.014$). There was no improvement by the combined use of H/E and MIB 1 slides. VIN 2 is far less confused with VIN 3 in the combined use of H/E and MIB 1 slides (9%) than in H/E slides (38%) ($p = 0.007$). There is a tendency to grade VIN in a two tailed grading system rather than a three tailed grading system, which became more apparent with the combined use of H/E and MIB 1 slides.

Conclusions—The interobserver variation with sole use of MIB 1 is better than with the use of H/E stain in VIN. The use of MIB 1 in grading VIN diminishes confusion between VIN 2 and VIN 3 fourfold. A two tailed grading system for VIN seems already to work in daily practice.

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Keywords: kappa test; MIB 1; vulvar intraepithelial neoplasia

Over 80% of patients with vulvar intraepithelial neoplasia (VIN) grade 3 present with multifocal disease. At microscopy, nearly 40% of these assumed multifocal lesions do not show VIN 3, but show VIN 2, VIN 1, or even histologically normal squamous epithelium.1

The advised international standard treatment for VIN 3 is surgical excision of all visible lesions, to exclude the presence of an occult invasive squamous cell carcinoma.2 However, a conservative approach in multifocal VIN 3, without histologically proven microinvasion, is also safe and effective.1 In this approach, invasive disease is excluded by taking multiple biopsies and the involved skin causing pain and pruritus is removed using cold knife surgery or laser vapourisation without aiming at radical removal.7 If one chooses to excise all visible lesions, it is therefore important to know which lesions do show VIN 3 and which do not. In this way one can leave as much vulvar skin as possible, avoiding psychological and sexual side effects from extensive surgery. It is not known how pathologists differ in their interpretation of VIN. However, they do differ in their interpretation of cervical intraepithelial neoplasia (CIN).1,4

Measurement of cell proliferation may provide useful information on diagnosis and tumour prognosis. The Ki-67 monoclonal antibody is currently used in evaluating cellular proliferation rates of malignant tumours.3 A formalin resistant epitope of Ki-67 cell proliferation associated antigen is immunohistochemically detected by the MIB 1 monoclonal antibody.5 This has been proven to be the best proliferation marker for routine use in formalin fixed and paraffin embedded tissue sections.6 In preinvasive and invasive squamous neoplasms of the uterine cervix, the number of MIB18 or Ki-67 positive cells increases with the severity of the lesion.10 Ki-67 expression has been described in normal vulvar epithelium and VIN 3,11 but its value has never been examined systematically in the grading of VIN. VIN can have a spectrum of pathological changes, such as nuclear pleomorphism, hyperchromasia, altered epithelial maturation, cellular crowding, loss of normal keratinocyte polarity, and atypical mitotic figures.11 VIN can have a spectrum of pathological changes, such as nuclear pleomorphism, hyperchromasia, altered epithelial maturation, cellular crowding, loss of normal keratinocyte polarity, and atypical mitotic figures.12 VIN has been subclassified as VIN 1, 2, or 3, based on the extent of cellular disarray of the epithelium.12 MIB 1 immunostaining is assessed by determining the percentage of labelled nuclei in the total population of nuclei and might therefore be a more accurate test than haematoxylin/eosin (H/E) staining for grading VIN.

In this study we sought to determine the interobserver variation in scoring the presence and grade of VIN in H/E slides, MIB 1 slides, and the combined use of H/E and MIB 1 slides.

Methods

Ten cases each of normal vulvar skin, VIN 1, VIN 2, and VIN 3 were retrieved from the pathology files of the department of pathology of the Academic Medical Centre. From each paraffin block two additional sections were recut. These sections were stained with H/E and MIB 1 (Immunotech, Coulter). The slides contained normal vulvar skin and VIN lesions. For MIB 1 antigen enhancement and optimisation of immunohistochemistry, slides were immersed in a 0.01 M sodium citrate dehydrated solution (pH 6.0) and then boiled for 10
minutes on a hot plate. After cooling overnight, the avidin-biotin complex methodology was used with diaminobenzidine as the chromagen. Nuclei were counterstained with haematoxylin. Five experienced pathologists and one gynaecological oncologist with special interest in gynaecological pathology each examined the 80 slides. They knew that their results would be compared, but they did not have any discussion over the grading criteria beforehand. Written definitions were given. First, the examiners scored the H/E and MIB 1 slides from each case separately, so that scoring of the H/E slides was not influenced by the corresponding MIB 1 slides and vice versa. On the H/E slides, the presence and grade of VIN (1–3) was scored according to the following definitions:\footnote{12}:

- Normal vulvar skin: no distorted architecture;
- VIN 1: cellular disarray involving the lower one third of the epithelium;
- VIN 2: cellular disarray involving the lower two thirds of the epithelium;
- VIN 3: cellular disarray involving more than the lower two thirds of the epithelium.

On the MIB 1 slides, the extent to which nuclei were positive throughout the epithelium was scored, depending on whether positive

![Figure 1 On the left from top to bottom: normal vulvar skin, VIN I, VIN II, VIN III. On the right from top to bottom: positive nuclei present in the basal two cell layers, the lower one third, the lower two thirds, and more than the lower two thirds of the epithelium.](image-url)
nuclei were present in either the basal two cell layers, the lower one third, the lower two thirds, or more than the lower two thirds of the epithelium. Outside the scored category for MIB 1, approximately 5% of positive nuclei in the remainder of the epithelium were considered as not present in the tissue sample. The kappa values for each pair of observers were calculated (table 3). The unweighted kappa values ranged from 0.22 to 0.63 for H/E slides, from 0.38 to 0.77 for MIB 1 slides, and from 0.41 to 0.77 for H/E–MIB 1 slides. The weighted kappa values ranged from 0.73 to 0.88 for H/E slides, from 0.85 to 0.95 for MIB 1 slides, and from 0.79 to 0.94 for H/E–MIB 1 slides. The kappa values were not better between pathologists than between pathologists and gynaecological oncologist, nor between pathologists within a single institute. The unweighted kappa was 0.47 for H/E slides, 0.62 for MIB 1 slides, and 0.60 for H/E–MIB 1 slides. The weighted kappa was 0.82 for H/E slides, 0.91 for MIB 1 slides, and 0.87 for H/E–MIB 1 slides. This improvement in unweighted and weighted kappa values between H/E slides and MIB 1 slides was significant (p = 0.023 and 0.014, respectively). However, the improvement in unweighted and weighted kappa values between H/E slides and H/E–MIB 1 slides was not significant (p = 0.08 and 0.26, respectively). In order to assess which in categories the H/E slides and the H/E–MIB 1 slides differed, unweighted and a weighted group kappa were calculated. We assumed that differences between group kappa values had a normal distribution, hence the significance of differences was assessed by calculation of z values.

**Results**

Of 720 observations (40 H/E slides, 40 MIB 1 slides, 40 H/E–MIB 1 slides, six observers), 719 were recorded, as observer 3 was unable to assign one score in a combined reading of H/E–MIB 1 slides. Tables 1 and 2 show the frequency distributions of the scores of the different categories for H/E slides, H/E–MIB 1 slides, and MIB 1 slides. In the MIB 1 slides, the presence of positive nuclei in the lower one third of the epithelium was scored less (8%) than in the other categories. This difference disappeared if the presence of positive nuclei in the basal two cell layers and the lower one third of the epithelium were considered together as one category (50%) and compared to the presence of positive nuclei in the lower two thirds and more than the lower two thirds of the epithelium, taken together as one category.

To compare the agreement between the individual observers for H/E slides, MIB 1 slides, and H/E–MIB 1 slides, unweighted and weighted kappa values for each pair of observers were calculated (table 3). The unweighted kappa values ranged from 0.22 to 0.63 for H/E slides, from 0.38 to 0.77 for MIB 1 slides, and from 0.41 to 0.77 for H/E–MIB 1 slides. The weighted kappa values ranged from 0.73 to 0.88 for H/E slides, from 0.85 to 0.95 for MIB 1 slides, and from 0.79 to 0.94 for H/E–MIB 1 slides. The kappa values were not better between pathologists than between pathologists and the gynaecological oncologist, nor between pathologists within a single institute. The unweighted kappa was 0.47 for H/E slides, 0.62 for MIB 1 slides, and 0.60 for H/E–MIB 1 slides. The weighted kappa was 0.82 for H/E slides, 0.91 for MIB 1 slides, and 0.87 for H/E–MIB 1 slides. This improvement in unweighted and weighted kappa values between H/E slides and MIB 1 slides was significant (p = 0.023 and 0.014, respectively). However, the improvement in unweighted and weighted kappa values between H/E slides and H/E–MIB 1 slides was not significant (p = 0.08 and 0.26, respectively).
the disagreements in grading are shown in table 4. The second column from the left (total number of scores) shows the combined frequency distribution of the scores of six observers for H/E slides and H/E–MIB 1 slides. The rows show the frequency distribution of the other five observers, when one observer assigned the score mentioned in the left hand column. For instance, for the slides that were scored as VIN 1 in the H/E slides by one observer, the distribution of the others were: 40% normal vulvar skin, 56% VIN 1, 4% VIN 2, and 0% VIN 3. Normal vulvar skin and VIN 1 were equally confused with each other (40–56%), but not with VIN 2 (4–9%) or VIN 3 (0%) in H/E slides. This sharp distinction between normal vulvar skin and VIN 1 on the one hand and VIN 2 and VIN 3 on the other hand appeared even more striking (though the difference was not significant) in the H/E–MIB 1 slides compared with the H/E slides, where no confusion was found at all. VIN 2 was hardly ever confused with normal vulvar skin, sometimes with VIN 1, but nearly twice as often with VIN 3 in H/E slides. VIN 2 is far less confused with VIN 3 in H/E–MIB 1 slides (9%) than in H/E slides (38%) (χ² test, p = 0.007).

Discussion

The κ statistic is the measure of choice for assessing interobserver variation. It corrects for chance agreement, which is not taken into account if percentages of agreement between observers are compared. Unweighted κ values depend on the number of categories one wants to distinguish: if detailed subdivisions are required, the task becomes more difficult and the κ values will be lower.11 This led to the development of the weighted κ,14 which take into account the degree to which disagreement concerns adjacent or more remote categories; κ values below 0.40 may be taken to represent poor agreement, values between 0.40 and 0.75 fair to good agreement, and values above 0.75 excellent agreement.15 Unweighted1 and weighted1 weighted κ values have been calculated for grading CIN on routinely stained H/E sections. It has been shown that the group κ for grading CIN is improved significantly if observers agree beforehand on which morphological characteristics should be considered relevant for grading CIN.13 This can be optimised after a consensus meeting between observers through a joint session behind a discussion microscope about the method of grading CIN.18 This may explain in part why in this study the interobserver variation for scoring presence and grading of VIN in H/E slides was already fair (unweighted group κ 0.47) to excellent (weighted group κ 0.82), as written definitions about grading were handed out at the scoring. Another explanation might be that the observers were all experienced in gynaecological pathology. Therefore, an improvement in unweighted and weighted group κ values between H/E slides and H/E–MIB 1 slides is difficult to establish. However, there was a significant improvement in unweighted as well as in weighted group κ values between H/E slides and MIB 1 slides. This is not surprising, as only one variable, namely positive nuclei, had to be taken into account in the MIB 1 slides. By contrast, many characteristics play a role in determining the presence and grading of VIN in H/E slides, such as the extent of cellular disarray of the epithelium, nuclear pleomorphism, hyperchromasia, altered epithelial maturation, cellular crowding, loss of normal keratinocyte polarity, and atypical mitotic figures. We assumed that the presence and extent of positive nuclei in four categories in the MIB 1 slides correspond with normal vulvar skin and VIN 1–3. It is unknown at present how MIB 1 immunoquantitation in general and the categories used in this study for presence and grade of VIN are correlated with clinical outcome of VIN. Therefore, the use of MIB 1 on its own for assessing the presence and grading of VIN is unwarranted at present. However, MIB 1 staining should be evaluated in correlation studies in which cytomorphometric analyses are also evaluated.

Positive nuclei present in the lower one third of the epithelium was scored significantly less in MIB 1 slides than in the other categories, although in H/E slides and H/E–MIB 1 slides such a strong difference was not found for VIN 1, compared with the other categories. This may be explained by the categories used for the presence of positive nuclei in MIB 1 slides in our study. The observers had to make a distinction between positive nuclei present in the basal two cell layers and the lower one third of the epithelium. In many VIN cases the basal two cell layers and the lower one third of the epithelium nearly enclose the same part of the epithelium. It is clear that the combination of H/E and MIB 1 significantly diminishes the confusion between VIN 2 and VIN 3, by 38% in the H/E slides and by 9% in the H/E–MIB 1 slides, as can be seen in table 4. It also seems from table 4 that observers were more likely to grade the slides into two categories (normal vulvar skin and VIN 1 or VIN 2 and VIN 3) than into four categories. We showed on the one hand that normal vulvar skin and VIN 1 are easily confused, but hardly ever with VIN 2 and never with VIN 3. This becomes even clearer in the H/E–MIB 1 slides. However, VIN 2 is hardly ever confused with normal vulvar skin, sometimes with VIN 1, but nearly twice as often with VIN 3. VIN 3 is never confused with normal vulvar skin, hardly ever with VIN 1, and sometimes with VIN 2. This finding of a two-tailed grading system for VIN may reflect the fact that CIN is graded as low grade squamous intraepithelial neoplasia (LGSIL) and high

Table 4  Frequency distribution (%) of the scores of five observers on 40 haematoxylin and eosin slides and 40 HE-MIB 1 slides (slides), conditional on the judgement of the other observer

<table>
<thead>
<tr>
<th>Vulvar skin Total number of scores (N)</th>
<th>Vulvar skin Normal (%)</th>
<th>VIN 1 (%)</th>
<th>VIN 2 (%)</th>
<th>VIN 3 (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal 54</td>
<td>47</td>
<td>69</td>
<td>44</td>
<td>31</td>
</tr>
<tr>
<td>VIN 1 62</td>
<td>53</td>
<td>40</td>
<td>48</td>
<td>56</td>
</tr>
<tr>
<td>VIN 2 48</td>
<td>59</td>
<td>4</td>
<td>9</td>
<td>22</td>
</tr>
<tr>
<td>VIN 3 76</td>
<td>73</td>
<td>0</td>
<td>0</td>
<td>3</td>
</tr>
</tbody>
</table>
grade squamous intraepithelial neoplasia (HGSIL). This could readily influence the grading of VIN.

CONCLUSIONS
Pathologists can use MIB 1 in grading VIN, as confusion of VIN 2 with VIN 3 is lowered fourfold. This could have implications for the management of patients with VIN 3, for whom standard treatment advice is to remove only VIN 3 lesions. It is worthwhile considering the implementation of a two tailed grading system—high grade and low grade squamous intraepithelial lesions—for VIN, as in daily practice this already seems to be what is done.

14 Cohen J. Weighted kappa: nominal scale agreement with provision for scaled disagreement or partial credit. Psychol Bull 1968;70:213–30.