Morphometric analysis of nuclei in epithelial structures from normal and neoplastic endometrium: a study using the Isaacs cell sampler and Endoscann instruments

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Summary Morphometric analysis of nuclear area and shape in epithelial cells from cytological specimens from 35 patients with normal endometrium and from 20 patients with moderately or well differentiated endometrial adenocarcinoma was performed. The mean nuclear area in malignant cells was significantly higher than in normal epithelial cells. The range of the mean values from the normal cases, however, included 70% of the malignant values. Furthermore, individual cell groups in a normal cell population often gave values well within the malignant range. The greatest distinction between normal and malignant cases was obtained using a cut off mean value of 45 \( \mu \text{m}^2 \). With this as the sole criterion 17% of reports would have been false positives and 25% false negatives.

Endometrial cytology has been in routine use in our region (Western Norway) since 1979–80, both in the gynaecological department at the university hospital and in gynaecological practice in the region. Accuracy tests on sampler material have shown that it may be as reliable as curettage and that cytological evaluation can be as accurate as histology in malignant cases, although there are reports of disappointing results.

It is commonly accepted that epithelial cell nuclei are smaller in normal than in malignant conditions of the endometrium. Evaluation of this sampler material, however, showed unexpected variation in nuclear appearance due mainly to variation in nuclear size. Smears in which the architecture of the glands was obviously normal on occasion showed pronounced nuclear enlargement, while smears with obvious malignant structure were often, on closer investigation, composed of almost normal looking cells. This study describes the variation in nuclear area and shape of epithelial cells seen in endometrial samples from a series of 35 normal and 20 malignant cases.

Material and methods

Thirty five cases with normal endometrial histology and 20 cases showing histologically moderately or well differentiated adenocarcinoma were studied. They were taken from a previous series of 400 patients in whom endometrial cell sampling using either the Isaacs cell sampler or the Endoscann instrument had been followed by curettage. Consecutive cases with the requisite diagnosis were selected from our files (Table 1). Two histopathologists had agreed on the histological diagnosis in each case.

When introduced into the uterine cavity both sampler instruments produced tissue fragments of surface epithelium, underlying stroma, and variable numbers of glands in normal cases. In malignant cases the material consisted of tumour tissue and variable amounts of the same material as in normal cases. For histological analysis the curettings had been fixed in 4% neutral buffered formaldehyde and stained with haematoxylin and eosin. The cytological material was smeared on to four glass slides, fixed in 96% alcohol, and stained by the Papanicolaou method. Only one of the four slides was used for measurement; the one used was the
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Table 1  Mean nuclear area (MNA) (±SD) in epithelial cells from 35 cytological samples from normal endometrium and 20 samples from patients with moderately or well differentiated adenocarcinoma of the endometrium

<table>
<thead>
<tr>
<th>Histological diagnosis (curettings)</th>
<th>Endoscann</th>
<th>Isaacs</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No of patients</td>
<td>MNA</td>
</tr>
<tr>
<td>Proliferation</td>
<td>5</td>
<td>40 ± 12</td>
</tr>
<tr>
<td>Secretion</td>
<td>5</td>
<td>43 ± 14</td>
</tr>
<tr>
<td>Atrophy</td>
<td>5</td>
<td>38 ± 9</td>
</tr>
<tr>
<td>Normal endometrium</td>
<td>15</td>
<td>40 ± 11</td>
</tr>
<tr>
<td>Adenocarcinoma (R)</td>
<td>10</td>
<td>62 ± 19</td>
</tr>
<tr>
<td>Adenocarcinoma (D)</td>
<td>10</td>
<td>67 ± 14</td>
</tr>
</tbody>
</table>

R = random measurement. D = diagnostic measurement.

first which on screening at ×25 magnification showed abundant material.

Measurements of nuclear area and nuclear shape were performed. The slides were evaluated with a light microscope (Leitz Dialux 20 EB) with a tracing device. A cursor with a light diode was placed on a digitising plate (Bit pad One TM Summagraphics Corporation, Fairfield, Connecticut) and with a drawing tube the light diode was projected on to the slide. The digitising plate was connected to a microcomputer (Commodore computer CBM 4032, Commodore Business Machines, St Clara, California). The nuclear circumference was followed with the light diode and a recording made at ×400

Fig. 1  Photomicrograph of a cytological smear from normal endometrium, ×225. (a) Typical example of a field measured in a superficial epithelial sheet. (b) Example of the monolayer chosen for measurement in glandular structures.
magnification. A morphometric program (MTS, Medizinisch Technische Apparate, Tübingen, West Germany) was used to measure the length of the circumference, the area of the nucleus (in \( \mu m^2 \)), and the shape of the nucleus, expressed as the form factor, simultaneously. A circle has the form factor 1.0.

In normal cases consecutive epithelial sheets or glandular structures were measured on routine screening of the slide. Twenty nuclei in each of 10 different epithelial structures were measured. The structures were excluded from measurement only if they did not contain at least 20 well preserved nuclei without artefacts such as stretching, crushing, or air drying. Pyknotic nuclei and mitotic figures were also excluded. Fig. 1 gives typical examples of a measurement field in a sheet of superficial epithelium and the monolayer field chosen for measurement in glandular structures. Measurement was done after optimal focusing of each nucleus.

In malignant samples two different types of recordings were made. One was performed in the same way as in normal samples, measuring nuclei in epithelial structures as they presented themselves on screening regardless of whether the observer judged them to be of normal or malignant origin. This was termed random measurement. In a second series of measurements on the same samples only clumps and sheets of epithelial cells which the observer diagnosed as malignant structures were measured; other structures were excluded. This type of measurement was termed diagnostic. Criteria for malignancy were the subjective assessment of architectural details such as papillary formations and irregular crowding of more or less well formed glandular formations, quality of the background material, cellular size, nuclear size, nuclear to cytoplasmatic ratio, nuclear staining...
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Properties, and nucleolar abnormalities.

Results

The mean nuclear area (MNA) in normal epithelial cells did not vary with hormonal state since an almost identical MNA was found in atrophic endometrium and in the proliferative and secretory phases of the normal cycle (Table 1). The MNA was also similar when measured on cells derived from the two samplers. The MNA for all measurements with both instruments for normal endometrium was $38 \pm 9 \mu m^2$. Table 1 also shows that malignant epithelial cells had a similar MNA on random and diagnostic measurement; the observed differences were not significant. There was no significant difference in the results from the Isaacs and the Endoscann instruments. The mean for all the malignant random measurements was $57 \pm 16 \mu m^2$ and for the diagnostic measurements $65 \pm 13 \mu m^2$. The differences in mean between normal material and malignant, $19 \mu m^2$ for random and $27 \mu m^2$ for diagnostic measurement, were both significant ($t$ test, $p < 0.001$ for both). Fig. 2 shows that the random measurements lay under the diagnostic in all but five cases. In these five cases the means were similar, the standard deviations overlapping.

Different cut off points were tested: a MNA of $45 \mu m^2$ (Fig. 3) proved to be the value that discriminated best between normal and malignant conditions ($\chi^2 = 18.1, p = 0.00002$). Only one case with atrophic endometrium showed values over $45 \mu m^2$. Three of 10 cases in proliferation and two of 10 in secretion exceeded this value. In contrast, 15 of 20

![Mean nuclear area for each of 55 samples related to standard deviation (SD).](image)

![Mean nuclear area from each cell group from each of 55 samples related to the condition of the endometrium. Each sample is presented as a column of points. Each point represents one or more cell groups.](image)
malignant cases were above and five below. If values over 45 μm² were considered indicative of malignancy there would have been six false positive (17%) and five false negative reports (25%).

Fig. 4 shows the relation between the MNA and its standard deviation in each of the 55 cases measured. For cases with MNA values below 45 μm² the standard deviation of the mean did not exceed 10 irrespective of whether the cases were normal or malignant. In contrast, when the MNA was over 45 μm² the standard deviation in about half the cases was over 10, again irrespective of whether the cases were normal or malignant. The scatter indicates great overlap in mean value in normal and malignant conditions. Only 17% of the MNA values obtained from malignant cases had readings above the upper level of normal conditions. Fig. 5 also indicates a wide range of MNA values both in the various normal conditions and in malignant cases. Further, an individual case with normal endometrium may show MNA values with a small range, while others show a wide range. The same holds for malignant cases. Cases with almost identical MNA values were found in malignant and atrophic endometrium.

Each of the points in Fig. 5 is the MNA from 20 nuclei. The mean of the variation coefficient for all these readings is related to the condition of the endometrium in Table 2. This variation, in contrast to that in Fig. 5, is an expression of the heterogeneity of the nuclei in the local cell population in a given endometrial condition. The figures show no difference between measurements with the two samplers: similarity in readings in normal conditions with slightly higher values in malignancy, irrespective of the type of reading, random or diagnostic. Thus the groups can probably be regarded as comparable.

In normal conditions the nuclei were round or nearly round (range 0.84–1.0). The range for malignant recordings was 0.65–1.0, but the mean value for recordings both in normal and malignant conditions was >0.9.

### Discussion

Modern cytological sampler material from the endometrium is a challenge to the cytologist in many ways. Firstly, one is unfamiliar with the type of material. The Isaacs cell sampler and Endoscann used in this study produce quite large tissue fragments in addition to sheets, small clumps, and single cells. Secondly, few systematic descriptions of the appearance of the cytological material from these samplers have been published and the diagnostic criteria have not been clearly defined. Reports that the mean nuclear area is greater in malignant epithelium,13–15 confirmed in this study, leaves one with the impression that malignant epithelium has larger nuclei than normal epithelium. This may be one explanation for the consistently, but not significantly, higher values for diagnostic compared with random readings found in the present study. Further, the general concept of variation in nuclear size and shape in malignant tissue leads one to expect a greater variability in carcinoma cases than in normal conditions.16 The present work shows that neither of these concepts holds in practice for well or moderately differentiated adenocarcinoma of the endometrium.

The combination of nuclear size, range in nuclear size, and shape of the nuclei is also of little help, since normal and malignant cases vary in the same way in these respects. A tendency to high MNA values combined with high standard deviations was seen in some of the malignant cases but was also found in benign samples. The low MNA values seen in some malignant cases are noteworthy. It is unlikely that the values represent measurement of normal structures in the samples, since curettings from these showed a clear predominance of tumour tissue.

From these results it follows that morphometric analysis of sampler material with a focus on nuclear size and shape cannot be used as a screening method for malignancy in the endometrium. It also follows that in subjective assessment of the material in routine diagnostic work one must not pay too much attention to nuclear size, but rather take other criteria into consideration, especially the architecture of the tissue and the quality of the background. As sampler material comes into more general use, the problems it presents will become increasingly important to both cytotechnologists and cytopathologists. They should be aware of the wide range of nuclear appearances in normal conditions of the endometrium. At the same time full awareness that well and moderately differentiated endometrial carcinomas may consist of cells in which the nuclei do not differ in size or shape from nuclei in normal cells is essential.

**Table 2 Mean of variation coefficient in normal endometrium and in moderately or well differentiated adenocarcinoma of the endometrium**

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<td>17 ± 4</td>
<td>17 ± 4</td>
</tr>
<tr>
<td>Secretion</td>
<td>16 ± 4</td>
<td>16 ± 4</td>
</tr>
<tr>
<td>Atrophy</td>
<td>17 ± 4</td>
<td>17 ± 5</td>
</tr>
<tr>
<td>Malignant (R)</td>
<td>19 ± 5</td>
<td>21 ± 5</td>
</tr>
<tr>
<td>Malignant (D)</td>
<td>21 ± 6</td>
<td>20 ± 5</td>
</tr>
</tbody>
</table>

R = random measurement; D = diagnostic measurement.
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References


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