A MORPHOLOGICAL PROTOCOL AND GUIDE-LIST ON UTERINE CERVIX CYTOLOGY ASSOCIATED TO PAPILLOMAVIRUS INFECTION

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SUMMARY

The present study was designed to further assess the validity of the cytological description of morphological lesions said to be related to Papillomavirus (HPV) infections in senior women. The casuistic comprised 196 cervical smears from a group of women with no clinical or morphological evidence of neoplasia, collected simultaneously with samples submitted to detection of HPV DNA by PCR in a previous study. Three experienced cytologists studied each slide in two different conditions, with an interval of 20 months between them. The first approach was performed under routine laboratory standards, whereas the second was guided by a list of 16 well-defined parameters indicative of HPV-related cytological lesions. When suspicious cases of HPV-related alterations were grouped with positive cases, they showed on average: sensitivity of 25.5%, specificity of 84.4% and positive predictive value (PPV) of 26.8%. When suspicious cases were grouped with negative cases, sensitivity decreased, whereas specificity and PPV increased, as expected. In the second reading, which followed a “guide-list”, a decrease in sensitivity was observed, contrasting with a sharp increase of positive predictive value. Among the 16 cytomorphological criteria tested, “koilocytosis”, “mild koilocytosis” and “condylomatous parabasal cells” yielded the best predictive value for HPV DNA detection by PCR. In conclusion, despite the low sensitivity, cytopathologic assessment of cervico-vaginal smears leads to a highly specific diagnosis of HPV infection in menopausal women, with PPV of 91.0% when directed by a guide-list of well-defined morphologic criteria.

KEYWORDS: HPV infection; PCR; Cervical smears; Cytomorphology.

INTRODUCTION

The morphological analysis of cervicovaginal cells by the Papanicolaou method is widely accepted as an effective screening test for cervical neoplasia. More recently, it has also been frequently employed to detect morphological changes related to human papillomavirus (HPV) infection. The causal relation between certain types of HPV, called “high-risk” types, and cervical cancer was established by several epidemiological and laboratorial studies. Non-productive (latent) infection by HPV is defined when viral replication only occurs in synchrony with the cellular cycle in normal squamous epithelium, and when the cytopathic effects related to HPV are not detected cytohistologically. The expression of the E4 viral protein in squamous epithelial cells, which produces a collapse in the cytokeratin matrix, possibly leads to the typical perinuclear caviation, which is one of the features of either clinic or sub-clinic HPV productive infection. On the other hand, the nuclear atypia related to HPV is due to the heteroploidy that appears as a result of fusiform mitotic abnormalities, which lead to the replication of the DNA without cytokinesis. The results of these interferences on the mitotic process are the formation of binucleate or multinucleate cells and larger atypical nuclei, followed by heteroploidy.

Sequences of viral genome, detected by molecular hybridization techniques, can remain in the host cell under episomal form or integrated to the DNA. Nuclear changes seem to be more frequent in samples in which integration of HPV DNA has occurred. When the Polymerase Chain Reaction (PCR) technique (which detects productive and nonproductive infection in normal cervical epithelial scrapes) is used, the frequency of infection by HPV in general population of sexually active women varies from 15 to 20%. Studies involving postmenopausal women are scanty, and the morphological alterations provoked by the HPV infection in the squamous cells are controversial. JOVANOVIC et al. (1995) reported the pseudo-koilocytic alterations in postmenopausal women, focusing on the possibility of false positive low-grade diagnosis under certain circumstances. The HPV detection in postmenopausal women is around 10%.

AYRE (1949) and PAPANICOLAOU (1954) were the first to document cytomorphological features which, later on, would be associated with HPV. KOSS & DURFEE (1956) defined “koilocytic atypia” as the presence of enlarged epithelial cells with irregular, hyperchromatic nuclei, encircled by transparent and clear space, from which the term “koilocytosis” derived. Infection by HPV is believed to
originate in the basal layer. After the replication of viral DNA in proliferative basal cells, the virus could, then, infect adjacent epithelial cells. Although the precise morphogenesis of lesions associated with HPV infection is not fully understood, possible cytopathic effects and organizational changes ascribed to HPV include: perinuclear cavity, binucleation, abnormal cytoplasmic keratinization, nuclear atypia and several degrees of nuclear degeneration. Recognition of these changes is subordinated to the quality of Pap smears collected, processing, triage and interpretation of microscopic images.

New cervical cancer screening strategies, such as cytology plus HPV DNA testing for women with cervical cytologic abnormalities, have been proposed. Our findings may contribute for the improvement of these strategies. The importance of studying this group is evident. Therefore, the aim of this study is to re-assess the value of cervical cytology to identify HPV infection in women who had previously undergone triage with cytological examination “negative for neoplasia”, comparing the performance under routine laboratorial conditions to the one directed by a pre-established guide-list of cytomorphological parameters. The presence of HPV DNA detected by PCR was considered the “gold standard”.

MATERIALS AND METHODS

Study population: The present investigation was carried out in cervicovaginal samples from women concomitantly submitted to a cytopathological study and to an investigation of HPV DNA by PCR. These women were selected as controls in a case-control study conducted in São Paulo City, Brazil, to investigate risk factors for invasive cervical cancer. Women between 18 and 79 years old (mean, 52; median, 51) were invited to participate, recruited from outpatient clinics and in-patient wards of five public hospitals in São Paulo City. The participants of the study had a pelvic examination performed by a gynecologist while collected.

Of the 225 controls of the previous study, 29 were excluded in the present study: 10 had no cytology taken, or the samples were considered unsatisfactory for interpretation; in 19 controls, the β-globin gene was not amplified, indicating that the DNA was inadequate for analysis by PCR. Remained, therefore, 196 women who had had cytological exam and PCR performed; 34 (17.3%) were positive for HPV DNA.

Cytological analysis: Three observers from the Institute Adolfo Lutz, the Central Public Health Laboratory of São Paulo State Health Secretariat, independently reviewed the original slides previously classified as “negative for neoplasia”.

Cytological interpretation comprised two reading stages. The first reading reproduced routine laboratorial conditions, conforming to the classification of the Brazilian Ministry of Health (1993) and the World Health Organization (WHO, 1994). Twenty months after the first reading, the same three observers performed a guided reading, following a pre-established guide-list with 16 cytomorphological parameters associated with infection by HPV, comprising “classic” and “non classic” parameters as listed (Table 1). Observers were not informed that they had previously analyzed those slides. In the first reading, 196 samples were evaluated; in the second reading, due to operational problems, 194 samples were evaluated.

<table>
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<tr>
<th>Marker</th>
<th>Defined by</th>
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<tr>
<td>Bi or multinucleation</td>
<td>Naib &amp; Masukawa, 1961; Meisels &amp; Fortin, 1976; Schneider et al., 1987</td>
</tr>
<tr>
<td>Karyorrhexis</td>
<td>Naib &amp; Masukawa, 1961; Takahashi, 1982</td>
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<td>Cleared cytoplasm</td>
<td>Schneider et al., 1987</td>
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<td>Spindle cells</td>
<td>Schneider et al., 1987</td>
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<td>Multinucleated giant cells</td>
<td>De Borges et al., 1989; Luzzatto et al., 1990</td>
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<td>Condylomatous parabasal cells</td>
<td>Meisels et al., 1988</td>
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<tr>
<td>Koilocytosis*</td>
<td>Koss &amp; Durfee, 1956; Meisels &amp; Fortin, 1976</td>
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<td>Condensation of filaments</td>
<td>Schneider et al., 1987</td>
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<td>Dyskeratocytes*</td>
<td>Meisels &amp; Fortin, 1976; Purola &amp; Savia, 1977</td>
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<td>Mild dyskeratosis</td>
<td>Schneider et al., 1987</td>
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<td>Perinuclear halos</td>
<td>Naib &amp; Masukawa, 1961; Meisels &amp; Fortin, 1976; Schneider et al., 1987</td>
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<td>Nuclear hyperchromatism</td>
<td>Meisels &amp; Fortin, 1976; Schneider et al., 1987</td>
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<td>Keratohyalin and keratohyalin-like granules</td>
<td>Schneider et al., 1987</td>
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<td>Smudge chromatin</td>
<td>Meisels et al., 1988</td>
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<td>Spindle nuclei</td>
<td>Shroyer et al., 1990</td>
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* “classic parameters”

Only in the second reading, observers filled in a specific form with all cytomorphological findings. At the end of the form, the exam was classified as positive, negative or suspicious for HPV infection. Observers were not aware of HPV DNA investigation by PCR.

Statistical analysis: Results of HPV DNA investigation by PCR performed at the Department of Pathology, Free University Hospital of Amsterdam, were considered reference (gold standard) to the presence or absence of infection by HPV. To evaluate the cytomorphological examination performance, sensitivity, specificity and positive predictive value (PPV) of the cytomorphological features associated with HPV were calculated with the respective 95% confidence interval (95% CI), using the Epi Info software.

In order to assess the validity of the classification “suspicious for HPV”, cases classified as such were analyzed twice: grouped together with the “positive for HPV” cases, and grouped with the “negative” cases.

RESULTS

First reading: Of the 196 samples, ten (5.1%) were considered inadequate by at least one observer and were excluded from the analysis. All the 34 positive samples for HPV DNA by PCR were among the 186 valid ones. In the first cytological reading of these 34 samples, under routine conditions, six (by observer A), two (by observer B) and three
(by observer C) were considered positive for HPV. Three, seven and five samples were classified as “suspicious for HPV” by observers A, B and C, respectively.

Among the 152 HPV DNA negative samples, 12 were considered positive by observer A, two by observer B and three by observer C. The three observers considered 26, 14 and 14 samples “suspicious”, respectively.

In Table 2, suspicious cases were grouped with positive ones. Such analysis showed on average: sensitivity of 26%, specificity of 84%, positive predictive value (PPV) of 27%. When suspicious cases were grouped with negative ones, as expected, sensitivity decreased, whereas specificity and PPV increased (Table 3).

**Second reading:** Of the 194 samples analyzed by the same three observers in the second reading, 16 (8.2%) were considered inadequate by at least one observer, remaining 178 valid samples. Since one of the inadequate samples corresponded to a woman diagnosed as infected by HPV by the PCR technique, 33 positive samples remained for the second reading.

The 33 positive cytological samples for HPV DNA by PCR were evaluated by the observers as follows: observer A - four positive and four suspicious for HPV; observer B - four positive and five suspicious; observer C - two positive and one suspicious.

In the evaluation of the 145 negative samples for HPV by the PCR technique, observer A considered one sample positive and three suspicious; observer B considered 16 samples suspicious, and observer C considered four samples suspicious. None of these samples were considered positive by observers B and C.

Overall, the performance of the cytological test was sharply improved comparing with the first reading. Such improvement was observed in both analyses - by grouping suspicious samples with either positive (Table 2) or negative ones (Table 3). Despite a small decrease in sensitivity, the second reading produced a large improvement in PPV. When the suspicious cases were grouped with those negative, PPV increased from 39% to 91% (Table 3).

Among the 16 cytomorphological parameters, “koilocytosis” presented the best PPV (mean 42%), followed by “mild koilocytosis” (mean 31%) and “condylomatous parabasal cells” (mean 28%) (Fig. 1).

| Table 2 |
|-----------------------|-----------------|-----------------|
| Observers (A,B,C) | First reading Mean (95% CI)** | Second reading Mean (95% CI)** |
| Sensitivity | 25.5 (12.7 - 44.0) | 20.2 (8.9 - 38.6) |
| Specificity | 84.4 (77.5 - 89.6) | 94.5 (89.4 - 97.6) |
| PPV * | 26.8 (13.4 - 45.9) | 45.5 (21.3 - 73.4) |

*PPV Positive predictive value; ** 95% Confidence Interval

**DISCUSSION**

In the present study, we assessed the value of cytology for the identification of HPV infection in a population of older women with cervical smears previously considered negative for neoplasia. These women participated as controls in a case-control study carried out in São Paulo City.

Koilocytic atypia defined by KOSS & DURFEE in 1956, and associated with HPV infection by MEISELS & FORTIN, 1976, has been the most reliable indicative of cytomorphological lesions related to HPV. Nonetheless, during the last decade, because of the demonstration of a relatively low sensitivity of cytology to identify this viral infection, several “non classic” parameters have been suggested, resulting in a considerable loss of specificity.

In the present study, in a casuistic of relatively old women, cervical cytology presented low sensitivity and high specificity for the identification of HPV infection detected by PCR. GJEN et al. obtained similar results in women without cervical dysplasia, aged between 20 and 44: sensitivity of 26% and specificity of 95%.
In the present investigation, sensitivity remained low, independently of either the way of grouping the women with results considered “suspicious for infection by HPV”, or the kind of reading (routine or guided) performed. However, specificity was higher when women with samples considered “suspicious” were grouped with those considered negative, and the reading was performed following a parameter guide-list. The greatest (and most important) difference between routine and guided readings was found in the positive predictive value, especially when “suspicious” samples were grouped with negative ones.

Considered the most sensitive method of detection of HPV infections, PCR detects productive as well as nonproductive infections. Many women with an HPV positive sample by PCR might be in a “nonproductive” stage, which explains the low sensitivity of cervical cytology. The interpretation of the cytomorphological features associated with HPV in Pap smears from postmenopausal women can be even more difficult. Since HPV needs epithelial differentiation to complete the development cycle, low levels of circulating estrogen reduce the maturation of squamous cervical cells. The high proportion of postmenopausal women may have also contributed to the low sensitivity found in this study.

The results obtained herein demonstrate that the reading following a parameter guide-list conducted to a marked improvement of the positive predictive value. They also show the inadequacy of using the terminology “suspicious changes of viral infection”. The sensitivity gains of such practice are of little significance, while reduction of specificity can bring undesirable worries to the patient and, occasionally, leading to inadequate therapeutic actions.

REFERENCES


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