Cervical cancer screening: from Pap smear to future strategies

Triagem de câncer do colo uterino: do teste de Papanicolaou a estratégias futuras

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ABSTRACT

Previously, the screening for detection of cervical cancer was performed by simple cervicovaginal sample collected by the physician whenever the patient attended the medical consultation, and soon it was established as the annual "Pap smear". Since then, an elementary test has evolved into a complex process with multiple algorithms for the identification of invasive disease. The detection of human papillomavirus (HPV) has become part of the new screening recommendations, resulting in major changes in the guidelines. This review intends to emphasize the most important topics that are part of cervical cancer screening, including cervical cytology and HPV detection, and to discuss particular aspects of cervical cancer in Brazil. Despite the great benefits achieved by the cervical cancer screening programs with cytology and HPV test, there are still important issues to be discussed and improved in defining future strategies, including simplicity and possible application in different socioeconomic contexts, definition of the best test or tests to be applied and recommended interval, minimizing possible harms. After the establishment of screening algorithms well defined by leading organizations, management protocols should be disseminated among physicians and patients by education programs.

Key words: cervical cancer; cervical cancer prevention; molecular diagnostic methods; vaginal smears; HPV DNA tests.

INTRODUCTION

Although the observation of cells was the first and original approach to the study of human diseases in the 19th century, the development of cytology as a diagnostic modality, as it is known today, followed the fundamental contribution of Dr. George Nicolas Papanicolaou, who first reported in 1928 that malignant cells from the cervix can be identified in vaginal smears. His work in collaboration with the gynecologist Herbet Traut provided a detailed description of the cytology of the female genital tract and the basis of the discovery of unsuspected occult cancer in asymptomatic patients, published many years later. Although initially their observations were received with skepticism by both pathologists and clinicians, many confirmed their findings subsequently, and cervical smears were embraced as a routine screening test for preinvasive lesions of the cervix, since then known as "Pap smear" or "Pap test"[1, 2].

The first cervical cancer screening clinics were established in the 1940s, when a number of women were screened for detection of early uterine cancer. In one year, it was found 54 cases of cancer in 639 women, 51 of them correctly diagnosed by cytology and six detected exclusively by Pap smear[3].

The implementation of a very simple and effective screening test was followed by a dramatic decrease in mortality rate related to cervical cancer in different populations[4-13].

Over the past 30 years, the widespread routine cervical cancer cytology screening has contributed to a 50% reduction in the incidence of cervical cancer in the United States. As demonstrated by the data, proper screening may effectively prevent cervical cancer, since 50% of women diagnosed with cervical cancer had never undergone cervical cytology testing and another 10% had not received screening in the five years preceding their diagnosis[14].

Cervical cancer is very rare among screened women[15].

Cervical cytology is reported according to the Bethesda system, which was introduced in 1988. The principles of the reports include clear, uniform, and reproducible terminology, reflecting the most current understanding of cervical neoplasia. It was revised in
1999, 2001 and the last updated version occurred in 2014, which includes an assessment of the specimen adequacy, whether there is evidence of lesions and the severity of the lesions(16).

**HUMAN PAPILLOMAVIRUS**

Most cervical cancers develop from infected cells with high-risk human papillomavirus (hrHPV), originated from the squamocolumnar junction. The causal link was described by Dr. Harald zur Hausen, who won the Nobel prize in 2008 for isolating the human papilloma virus (HPV) types 16 and 18 from cervical cancer tissue(17).

HPV is among the most powerful human carcinogens and has been implicated not only to cervical cancer, but also to cancers at several sites. HPV infection is the most common sexually transmitted infection worldwide, mainly in low- and middle-income countries(18). Apparently hrHPV is a necessary but not a sufficient condition for almost all cervical cancers. The risk of preinvasive lesions and invasive cancer of the cervix is strongly associated with persistent infection with hrHPV, especially type 16. Fortunately, most HPV infections in human are harmless, and cause no lesion. Due to the interaction between host and the pathogen, the majority of infected women will clear the virus and the precancerous lesions will regress. Only about 1% of low-grade lesions (CIN1) and 12% of high-grade lesions (CIN3) will progress and become invasive if left untreated(19).

Since the causal link between cervical cancer and hrHPV infection was established, much effort has been devoted to the study of prevention and identification of HPV infection. Currently vaccination against carcinogenic strains of HPV is commercially available, but even in some developed countries the vaccination uptake has been slow(19, 20).

**PAPANICOLAOU TEST – “PAP SMEAR”**

In use for more than 50 years, the Pap test in its original preparation, also called “conventional cervical smear (CS) method for cytology collection”, is still acceptable for screening purposes(21). It remains as an alternative for cervical cancer screening due to its simplicity and low cost.

The liquid based cytology (LBC) was approved by the US Food and Drug Administration (FDA) in 1996 as an alternative to conventional cervicovaginal smear. Although several studies showed an increased detection of low- and high-grade squamous intraepithelial lesions (LSIL and HSIL) by LBC preparations, a systematic review did not confirm that LBC is more accurate than conventional smears, but has an equivalent performance. Even so, LBC is gradually replacing the conventional cytological preparations, because it presents many other clear advantages, including the possibility of aliquoting for the hrHPV test(22).

Cervicovaginal cytology is clearly far from being a perfect screening test. In a systematic review, it was shown to have a sensitivity of only 51% (ranging from 30% to 87%) and a specificity of 98% (ranging from 86% to 100%), although methodological quality and frequency of histological abnormalities varied greatly; and only 12 of the 94 studies with less biased estimates were analyzed(23).

Furthermore, there is a significant interobserver variability in the interpretation of cytology, contributing to variations in sensitivity and specificity rates. We must not forget that the interpretative variability is also significant for histological specimens, even among well-trained observers demonstrated in cervical biopsies, as well as many other sites and organs(24).

A meta-analysis found that about 29% of failures to avoid invasive cervical cancer can be attributed to false-negative cytology. The authors examined 42 studies from 1950 to 2007, and the most common failure of the process was history of poor screening: 54% of women had inadequate screening intervals and 42% had never been screened. It should be emphasized that the proportion of Pap smears originally reported as normal and that, after review, were classified as false-negatives, or those normal cases that were not reviewed, but simply assumed to be false-negative, varied greatly among studies. Sampling errors may have contributed to at least some of the cases not reviewed(25).

**HPV TEST**

The detection of HPV deoxyribonucleic acid (DNA) may be accomplished by several molecular methods, particularly including signal-amplification (Hybrid Capture® assay) and polymerase chain reaction (PCR) based methods. The Hybrid Capture® system is designed to detect HPV divided into high- and low-risk groups, without genotyping individual virus, demonstrating high sensitivity and specificity for hrHPV. The PCR-based techniques are highly sensitive and specific but, besides being a labor-intensive procedure, it also presents some drawbacks, such as false negative results. Real-time PCR assay is a rapid, reproducible and reliable diagnostic tool, which has the additional advantages of detecting very small viral concentrations and different targets simultaneously, as well as to determine the viral load(26).
Initially, the hrHPV test was recommended as a reflex testing of atypical squamous cells of undetermined significance (ASC-US) to screen patients to colposcopy. The sensitivity for detecting CIN3 or greater by hrHPV test was 96.3% compared to 44.1% of cytology by one repeat with a screening threshold of HSIL or greater\textsuperscript{(27)}.

In 2004, the National Institute of Health, the National Cancer Institute, the American Society for Colposcopy and Cervical Pathology (ASCCP) and the American Cancer Society (ACS) agreed to expand the use of hrHPV as a cotesting\textsuperscript{(28, 29)}.

The joint recommendation released in 2012 advocates HPV testing to be used in conjunction with routine cytology, as well as a reflex testing in women aged 30 to 65 years\textsuperscript{(14, 21)}. A woman with a negative result for both hrHPV and cytology has a lower risk of developing CIN2 or CIN3 in the next four to six years\textsuperscript{(14)}. The cotesting increases the detection of CIN2 or greater lesions at baseline and significantly decreases the detection rates of CIN2/ CIN3 or greater lesions at subsequent screening compared to cytology alone\textsuperscript{(30)}.

The ATHENA trial demonstrated that 10% of women who tested positive for HPV 16 and/or 18 had high-grade cervical neoplasia (CIN2 or worse) that was not detected by cytology\textsuperscript{(31)}. Many other studies have documented that HPV testing has a greater sensitivity and reproducibility with increased negative predictive values compared to cytology\textsuperscript{(32-38)}. The data presented by these studies endorsed the idea of hrHPV test as a primary screening test, replacing cytology as screening women for colposcopy, advocated by many.

In 2014, the FDA approved the cobas\textsuperscript{®} HPV test for primary screening of cervical cancer in women aged 25 or older, by detection of hrHPV genotypes, at intervals equal to or greater than three years\textsuperscript{(39)}.

Intermediate guidelines for primary hrHPV screening were developed by representatives from the Society of Gynecologic Oncology, the American Society of Cytology, and the College of American Pathologists, in addition to American Congress of Obstetricians and Gynecologists (ACOG), and all groups authoring the 2012 screening guidelines\textsuperscript{(40)}.

In the Quest Diagnostics Health Trend study, which enrolled more than 250,000 women, the HPV/Pap cotesting identified more women whose cervical biopsy result revealed a finding of CIN3 or greater than HPV-only testing (98.8% versus 94%). The data showed that up to 19% of women with cancer may be missed when they are screened using HPV testing only; HPV/Pap cotesting misses 5.5% of cancer cases, and Pap alone, 12.2%. The authors concluded that cotesting in women aged 30-65 is the most effective screening test for detecting cervical cancer\textsuperscript{(41)}.

Although the cobas\textsuperscript{®} HPV test has been approved, it was stressed that it has limited sensitivity (27% for women equal to or older than 50 years, 36% for women equal to or older than 40 years, 53% for women equal to or older than 30 years, and 58% for women equal to or older than 25 years), which is much lower than that observed with the cotesting in women equal to or older than 30 years, perhaps because of suboptimal performance of cytology\textsuperscript{(31, 39, 42)}. Negative HPV rates in patients with invasive cervical cancer varied among authors and seemed to increase with time before cancer diagnosis, perhaps due to the smaller size of lesions or lower viral titers during earlier periods\textsuperscript{(42)}. Another arguable point is the use of CIN2 or/and CIN3 or worse as the right endpoint for evaluating cervical screening algorithm, since it does not reflect cancer risk accurately\textsuperscript{(31, 42)}.

**SCREENING GUIDELINES**

Screening recommendations proposed by several societies and private organizations have been published and are reviewed periodically when new evidence suggests that a change may be necessary. Previous established guidelines showed considerable variation prior to 2012.

The current American guidelines for cervical cancer screening were created in 2012 as a joint recommendation of the ACS, ASCCP and the American Society for Clinical Pathology (ASCP), which were accepted and promoted by the ACOG, in association with US Preventive Service Task Force (USPSTF). The benefit was defined as more detection of CIN3 or worse at baseline and reduction in CIN3 or worse detection at subsequent rounds of screening. The harm was defined as an increase in number of colposcopy\textsuperscript{(14, 21, 40)}.

Some reviews on cervical cancer screening guidelines present a summary from major organizations recommendations and the reader will find more detailed information\textsuperscript{(44)}.

The current main guidelines recommendations are the following:

- cervical cancer screening should begin at 21 years of age, regardless of age of coitarche or vaccination status, with cervical cytology tests exclusively until 30 years of age (either with conventional or liquid-based cytology), every three years;
- for women from 30 to 65 years of age, cotesting with cytology and HPV test every five years is preferred, although cytology screening every three years is acceptable;
- screening should be discontinued for women over 65 years of age at low risk, with no history of cervical intraepithelial neoplasia (CIN) grade 2 or greater, with negative results in prior screening;
screening should be discontinued for women of any age who have total hysterectomy and have no history of cervical cancer or precancerous condition.

Adherence to guideline recommendations is quite variable. Many clinicians continued to suggest annual Paps, as recommended by ACOG, although current guidelines advocated against annual screening, since no advantage is observed in relation to Pap tests performed every two or three years. Physicians believe that patients were uncomfortable with less frequent testing and if they extend the screening intervals, patients would not return annually just for the clinical examination(45).

CERVICAL CANCER IN BRAZIL

According to the 2016/2017 estimates, Brazil will register next year 300,800 cases of cancer among women(46). More than 16,000 new cases of cervical cancer are expected in 2016(47). Cervical cancer remains as the third leading cause of cancer-related mortality among women for decades, without any improvement(48).

Currently, the Brazilian program to cervical cancer control is based on population screening and vaccination, used together as complementary actions and coordinated by the Brazilian National Cancer Institute (Instituto Nacional de Câncer José Alencar Gomes da Silva [Inca]), an agency of the Ministry of Health of Brazil facing national integrated actions for the control and prevention of cancer(49). The screening method is the Pap test or Pap smear for women between 25 to 64 years old, or sexually active women; this test is provided annually (or once every three years after two normal tests) and it is followed by colposcopy for HSIL, carcinoma, or persistent LSIL or ASC-US.

LBC was also incorporated as the standard method of evaluating cervical samples in Brazil, largely replacing CS. Some Brazilian studies have demonstrated a better performance of LBC compared to CS, with lower rates of unsatisfactory specimens and higher sensitivity(50-52). A more recent study critically analysed 218,594 cases collected in a public health service in the state of São Paulo and observed positivity of 5.7% versus 3%, respectively, in LBC and CS; unsatisfactory preparations were present in 0.3% and 3% of the cases, respectively(53). However other groups have observed similar performances between the methods, finding no significant differences(54, 55).

In 2014, the Ministry of Health of Brazil launched the National Immunization Program through a quadrivalent HPV vaccine (subtypes 6, 11, 16, and 18) for girls between 9-13 years old(57).

A recent study evaluated the cervical cancer screening program in Brazil from 2006 to 2013 using the Information System of Cervical Cancer Screening (Sistema de Informação do Câncer de Colo de Útero [SISCOLO]), created by the Department of Informatics of the Public Health System (Departamento de Informática do Sistema Único de Saúde [DATASUS]), which contains information on all Pap tests collected in the public health system, and was implemented for the management and monitoring of the cervical cancer screening program(58). A decreasing trend in the rates of LSIL and HSIL was observed, as well as lower numbers of positive cytological diagnosis and an increased rate of rejected exams. The positivity rates and the frequency of unsatisfactory cases were lower than expected. The authors suggest that actions should be taken by the government to improve the effectiveness of cervical cancer control in Brazil, through more funding for internal quality control during both the pre-analytical and the analytical phase(59).

Albeit Brazil, like many other countries in Latin America, has a cytology-based screening program, they often have problems with quality and/or delays in follow-up care(60).

FUTURE STRATEGIES

The best screening algorithm remains a matter of debate.

Primary HPV screening is an attractive option to health service because the results are not subject to inter-observer variation. However, it requires equipments, reagents, personnel, training, quality control and accreditation. This scenario is far from the real world in different populations, even in developed countries, considering that many women will be screened or may never be screened at all.

We must remember the fact that the system for cervical cancer screening with both the Pap test and the HPV test is already working in many practices. There is no reason to disrupt such an operative scheme that is working successfully without adequate evidence of additional benefit of primary HPV screening. Further data are needed on the actual benefits and costs and the impact on the use of colposcopy and other diagnostic tests(61).

Supporters of primary HPV screening claim that this method not only finds more CIN3 or worse than cytology or cotesting does, but also find them earlier; moreover, the positive predictive value of the primary HPV screening algorithm was greater than that
one of cytology. In contrast, those who advocate cotesting believe this approach detects more disease than the HPV test alone and emphasize that the performance of this new algorithm has not been assessed in routine clinical use. The focus of the debate about the best screening algorithm to detect cervical cancer is much more complex, since it may involve reducing costs rather than maximizing protection, not only by decreasing the number of tests, but also by increasing the screening intervals.

In some practices, where access to cytological examination is limited, primary HPV testing may allow to provide screening for the patients, which had not been previously possible.

Several studies support that HPV testing is feasible in low-resource setting as a tool for cervical cancer screening. The incorporation of new technologies, adapted for low- and middle-income countries may be part of future programs for early diagnosis and control of the disease. Although the best screening strategy in this context is still a work in progress, perhaps HPV testing can be applied using a self-obtained vaginal samples that will allow first-line screening and triaging of HPV-positive women during a single visit, defining management and eventually treatment.

In summary, despite the great benefit that the cervical cancer screening programs achieved through the use of cytology and HPV testing, there are still important issues to be discussed and improved in defining future strategies, including simplification and possible application in different socioeconomic contexts, definition of the best test or tests to be applied and interval recommendation, minimizing harms.

After well defined, screening algorithms were established by leading organizations; management protocols should be disclosed among physicians and patients through education programs, integrated into a multidisciplinary team, with the participation of all professionals involved in women’s health, ensuring not only a more effective diagnosis, but also an appropriate treatment and monitoring, connecting the primary, secondary and tertiary levels of health. In Brazil, the new recommendations are being finalized and will soon be published by Inca, Ministry of Health of Brazil.

CONCLUSION

1) Cervical cytology, including conventional smear and liquid based cytology, is a successful method for cancer screening and is still recommended as the exclusive test for women 21 to 29 years of age; 2) since HPV was established as the main causative agent of cervical cancer, its detection improved screening sensitivity; 3) cotesting, hrHPV test used in association with cytology, is recommended for women 30 to 64 years of age, since it is the most effective screening method for detecting cervical cancer; 4) the sole routine clinical use of HPV testing, or primary HPV testing, is still a matter of debate, but, perhaps, it may prove to be an option for strategic screening in countries with limited resources, as new tests are becoming faster, automated and cost-effective.

RESUMO

Inicialmente, a triagem para detecção do câncer de colo uterino era feita por meio de uma simples amostra cervicovaginal coberta pelo médico, sempre que o paciente comparecia à consulta médica; logo se estabeleceu como “exame de Papanicolaou” anual. Desde então, um teste elementar evoluiu para um processo complexo, com múltiplos algoritmos para identificação de doença invasiva. A detecção do papilomavírus humano (HPV) tornou-se parte das novas recomendações de triagem, resultando em grandes mudanças nas diretrizes. Esta revisão pretende enfatizar os tópicos mais importantes que fazem parte do rastreamento do câncer de colo do útero, incluindo citologia cervical e detecção do HPV, bem como discutir aspectos particulares do câncer de colo do útero no Brasil. Apesar dos grandes benefícios alcançados pelos programas de rastreamento do câncer de colo uterino por meio do uso da citologia e do teste do HPV, existem ainda pontos importantes a serem discutidos e melhorados na definição de estratégias futuras, como simplicidade e possível aplicação em diferentes contextos socioeconômicos, definição do melhor teste ou testes a serem aplicados e intervalo recomendável, minimizando possíveis danos. Após o estabelecimento de algoritmos de rastreamento bem definidos pelas principais organizações, protocolos de manejo devem ser divulgados entre médicos e pacientes por programas de educação.

Unitermos: neoplasias do colo do útero; prevenção de câncer de colo uterino; técnicas de diagnóstico molecular; esfregaço vaginal; testes de DNA para HPV.
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