Effective screening programmes for cervical cancer in low- and middle-income developing countries

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Abstract Cervical cancer is an important public health problem among adult women in developing countries in South and Central America, sub-Saharan Africa, and south and south-east Asia. Frequently repeated cytology screening programmes — either organized or opportunistic — have led to a large decline in cervical cancer incidence and mortality in developed countries. In contrast, cervical cancer remains largely uncontrolled in high-risk developing countries because of ineffective or no screening. This article briefly reviews the experience from existing screening and research initiatives in developing countries.

Substantial costs are involved in providing the infrastructure, manpower, consumables, follow-up and surveillance for both organized and opportunistic screening programmes for cervical cancer. Owing to their limited health care resources, developing countries cannot afford the models of frequently repeated screening of women over a wide age range that are used in developed countries. Many low-income developing countries, including most in sub-Saharan Africa, have neither the resources nor the capacity for their health services to organize and sustain any kind of screening programme. Middle-income developing countries, which currently provide inefficient screening, should reorganize their programmes in the light of experiences from other countries and lessons from their past failures. Middle-income countries intending to organize a new screening programme should start first in a limited geographical area, before considering any expansion. It is also more realistic and effective to target the screening on high-risk women once or twice in their lifetime using a highly sensitive test, with an emphasis on high coverage (>80%) of the targeted population.

Efforts to organize an effective screening programme in these developing countries will have to find adequate financial resources, develop the infrastructure, train the needed manpower, and elaborate surveillance mechanisms for screening, investigating, treating, and following up the targeted women. The findings from the large body of research on various screening approaches carried out in developing countries and from the available managerial guidelines should be taken into account when reorganizing existing programmes and when considering new screening initiatives.

Keywords Cervix neoplasms/diagnosis/prevention and control; Cervix uteri/cytology; Vaginal smears/utilization; Mass screening/organization and administration; Developing countries; Central America; South America; Africa South of the Saharan; South Africa; India; South-East Asia (source: MeSH).

Mots clés Tumeur col utérus/diagnostic/prévention et contrôle; Col utérin/cytologie; Frottis vaginal/utilisation; Dépistage systématique/organisation et administration; Pays en développement; Amérique centrale; Amérique du Sud; Afrique sub-saharienne; Afrique du Sud; Inde; Asie Sud-Est (source: INSERM).

Palabras clave Neoplasmas del cuello uterino/diagnóstico/prevencción y control; Cuello uterino/citología; Frotis vaginal/utilización; Tamizaje masivo/organización y administración; Países en desarrollo; América Central; América del Sur; África del Sur del Sahara; Sudáfrica; India; Asia Sudoriental (fuente: BIREME).


Voir page 960 le résumé en français. En la página 961 figura un resumen en español.

Introduction Cervical cancer is an important public health problem for adult women in developing countries in South

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and Central America, sub-Saharan Africa, and south and south-east Asia, where it is the most or second most common cancer among women. The vast majority of cervical cancer cases are caused by infection with certain subtypes of human papilloma virus (HPV), a sexually transmitted virus that infects cells and may result in precancerous lesions and invasive cancer (1). Developing countries accounted for 370 000 out of a total of 466 000 cases of cervical cancer that were estimated to occur in the world in the year 2000 (2). Worldwide, cervical cancer claims the lives of 231 000 women annually, over 80% of whom live in developing countries. A conservative
estimate of the global prevalence (based on the number of patients still alive 5 years after the diagnosis) suggests that each year there are 1.4 million cases of clinically recognized cervical cancer. It is also likely that 3–7 million women worldwide may have high grade dysplasia.

Some of the developing countries that have data on cancer incidence and/or mortality have registered either a stable or slowly declining trend in cervical cancer incidence, most likely due to sociodemographic changes rather than to early detection/prevention efforts (3). On the other hand, some regions in sub-Saharan Africa have registered an increased incidence in recent years (4). Despite the declining trends in incidence observed in some regions, the total burden of cervical cancer is rising in high-risk developing countries, mostly due to increasing populations.

In developed countries, initiation and sustenance of cervical cytology programmes involving the screening of sexually active women annually, or once in every 2–5 years, have resulted in a large decline in cervical cancer incidence and mortality (Fig. 1 and Fig. 2) over the last 40–50 years (5–8). The aim of these programmes is to detect precancerous lesions and treat them before they progress to invasive cancer. In contrast, the risks of disease and death from such lesions have remained largely uncontrolled in high-risk developing countries, mostly because of the lack of screening programmes or because of their ineffectiveness. This paper reviews existing experiences, achievements, constraints, and lessons learned in community-based, cervical cancer intervention programmes in developing countries. The sensitivity and specificity values that we report for various screening tests correspond to the detection of high-grade lesions (cervical intraepithelial neoplasia II and III) and invasive cancer.

Cervical cytology screening programmes worldwide

To date, cervical cancer prevention efforts worldwide have focused on screening sexually active women using cytology smears and treating precancerous lesions. It has been widely believed that invasive cervical cancer develops from dysplastic precursor lesions, progressing steadily from mild to moderate to severe dysplasia, then to carcinoma in situ, and finally to cancer. It now appears that the direct precursor of cervical cancer is high-grade dysplasia, which in about a third of instances may progress to cervical cancer over a period of 10–15 years, while most low-grade dysplasias regress spontaneously (9, 10).

Even though the impact of cytology screening has never been proved through randomized trials, it has been shown to be effective in reducing the incidence and mortality from cervical cancer in developed countries (5–8). The incidence of cervical cancer can be reduced by as much as 80% if the quality, coverage, and follow-up of screening are high. In most developed countries, women are advised to have their first smear test soon after becoming sexually active and subsequently once every 1–5 years. Many national guidelines are currently moving towards less frequent smear tests (once every 3–5 years) because it is recognized that cervical lesions develop slowly over several years. Women with low-grade lesions are generally advised to return for routine follow-up smears. Women with high-grade precursor lesions are further evaluated via colposcopy, biopsy, and subsequent treatment of confirmed lesions. Organized programmes with systematic call, recall, follow-up and surveillance systems that have shown the greatest effect (e.g. in Finland and Iceland), even though they use fewer resources than unorganized programmes (e.g. in the USA).

Cervical cytology is considered to be a very specific test for high-grade precancerous lesions or cancer but, even if the quality of collection and spreading of cells, fixation, and staining of smears, and reporting by well-trained technicians and cytopathologists are good, its sensitivity is only moderate. The results of meta-analyses suggest that cytopathological screening has a very wide range of sensitivity to detect lesions (11, 12); for example, cytology is estimated to have a mean sensitivity of 58% and a mean specificity of 69% (11). Also, estimates of sensitivity of conventional cytology (for high-grade lesions) vary greatly in individual studies, by as much as 30–87% (mean, 47%) (12). Both sampling and detection (reading) errors probably contribute to the low-to-moderate sensitivity of cytology. Assuming that cytology is only moderately sensitive, it seems likely that the observed decline in the risk of cervical cancer in developed countries may have arisen from the high screening frequency. Cervical neoplasia is a disease that progresses slowly, and many low-grade precancerous lesions regress spontaneously or do not progress further. High-grade lesions that are missed in a given screening round would probably be detected during the subsequent rounds in a frequently repeated cytopathological screening programme. A critical review of conventional cervical cytology in developed countries, where it was shown to be effective in cervical cancer control, provides valuable leads for public health policy decisions in low-resource environments. Current procedures, involving screening women once every 1–5 years, have considerable cost and resource implications. The limited health care budgets in most developing countries preclude initiating and sustaining such programmes, even in a limited geographic setting.

Cervical cancer screening programmes in developing countries

Cytology-based screening programmes for cervical cancer have been introduced in some developing countries, particularly in South and Central America,
over the last 30 years, but generally have achieved very limited success. In contrast, a comparison of the performance of conventional cytology and its potential alternatives in detecting cervical cancer and its precursors is ongoing in Asia, Africa, and Latin America. Both these approaches are briefly reviewed below since they provide potentially useful information for directing public health policy on introducing new and effective programmes in low-resource settings and for reorganizing existing programmes.

South and Central America

Since the 1970s there have been efforts to organize cervical cytology screening programmes nationally or regionally in selected Latin American countries. Chile. In Chile the cervical cancer screening programme has been in operation since the early 1970s. Cervical cancer mortality rates did not change much in the period from 1970 to 1985 after the introduction of the programme (Fig. 2). A recent evaluation of the programme indicated that more than 80% of the married women in Chile have been screened at least once. The programme was reorganized in the early 1990s, and mortality from cervical cancer has subsequently begun to decline.

Colombia. In Colombia, the Colombian National League against Cancer (a part of the public health system) and private organizations such as PROFAMILIA have been offering cytology screening since the 1970s (13). Subsequently, the cervical cancer mortality rate in the country has, however, remained stable (Fig. 2). Nevertheless, there has been a steady and substantial decline in the incidence of cervical cancer in the city of Cali (Fig. 1), possibly as a result of the ongoing screening activities carried out there since 1967, including educational and early detection campaigns. In 1990, a 5-year nationwide cervical cancer control programme was initiated to provide cytology smears to more than 60% of women aged 25–69 years over a 3-year period and to provide follow-up to over 90% of the women screened. The programme trained over 4000 nurses, 40 gynaecologists, and 36 pathologists. Cytology services were centralized and extensive community information and education campaigns were launched. Midway through the project, the centralized national health care system was reorganized and several services were decentralized to encourage the creation of efficient networks of services and surveillance. However, 5 years after the initiation of the programme, cervical cancer mortality data suggested that the situation had remained unchanged.

Costa Rica. In Costa Rica nationwide cytology services have been available to women aged ≥ 15 years since 1970. Information/education campaigns have been used to encourage sexually active women to have annual cytology smears. Invariably in all pelvic examinations a Pap smear is also obtained (14). Annually, around 250 000 smears have been performed and reported. Coverage has varied considerably according to region, with coverage of rural areas being inadequate in each given round of screening. Despite this, it seems that more than 85% of eligible women have been screened at least once. Though cervical cancer incidence remained virtually unchanged from 1983 to 1991, a significant decline has been observed more recently (i.e. a 3.6% decrease in annual incidence in 1993–97 compared with that in 1988–92) (Fig.1). However, the cervical cancer mortality rates have remained unchanged over the last 25 years (Fig. 2) (15, 16). In an ongoing cohort study of more than 9000 women in Guanacaste Province, the cross-sectional sensitivity of HPV testing was found to be higher than that of conventional cytology (88% versus 78%) but the specificity was lower (89% versus 94%) (17).

Cuba. In Cuba a cervical cytology screening programme, offering smears every two years to women aged ≥ 20 years, was implemented through...
the primary health care services in 1968 (18). Pap
smears are taken by a nurse in the family doctor’s
office and are processed in one of the 36 regional
cytology laboratories. It has been suggested that
more than 80% of Cuban women aged 20–60 years
have been screened at least once. However, no
reduction in cervical cancer incidence and mortality
(Fig. 2) has been observed since the introduction of
the programme.

**Mexico.** A national cervical cancer screening
programme was initiated in Mexico in 1974 (19, 20)
and now operates in the Federal District and all
31 states of the country. Cytology smears are offered
annually to women aged 25–65 years and the
programme is integrated with the existing health
care services. Mexico reportedly had 463 cyto-
technologists, 251 reading centres, 70 dysplasia
clinics, and 540 gynaecological oncology units in
1996. However, the infrastructure and resources
were sufficient to carry out only 3.5 million smears
annually for a target population of 16.5 million
women (data for 1996); annual screening was
nevertheless the “norm” for the programme.
Realistically, this infrastructure is sufficient to screen
the targeted women only once every 5 years. The
Ministry of Health (MOH) has a total of 120 cervical
cancer screening centres (CCSCs) where 230 cyto-
technologists are employed. These screening centres
are intended to carry out cytology screening of
6.5 million women who are not covered by social
security. The Mexican Institute for Social Security
(IMSS) is responsible for screening women covered
by social security. In 1992, the MOH’s screening
centres carried out 1.02 million smears and the IMSS
1.3 million smears. There is a wide variation in the
coverage of women on the national level. Studies
indicate that less than 30% of the women in rural
areas have been screened so far. There is no
systematic effort to coordinate the programme through a
central organization for call, recall, and follow-up of screened women.

An evaluation of the cervical cytology tests
provided within the Mexican programme indicated
that the validity and reproducibility varied greatly
within and between the screening carried out by the
MOH and the IMSS (21). Among the CCSCs
the sensitivity to detect high-grade lesions varied
from 46% to 90% and that of the specificity from
48% to 96%. The false-negative rate varied from
10% to 54%, with an average false-negative rate of
35%. Review of a random sample of 6011 nega-
tive smears indicated that 64.0% of the smears
were of insufficient quality. There has been no
decline in mortality from cervical cancer in Mexico
since the initiation of the screening programme
(Fig. 2) (22).

**Brazil, Peru, and Puerto Rico.** There are no
organized cervical cancer screening programmes in
Brazil. A high-risk of the disease (incidence >40 per
100 000 women) is reported from the north-east
region. Low-level sporadic screening with opportu-
nistic cytology smears is carried out in different
regions.

Peru has also recorded a high incidence of
cervical cancer; there are no organized screening
programmes in the country. A large demonstration
project of cervical cancer screening with visual
inspection with acetic acid (VIA) is currently on-
going in San Martin region of Peru.

An early detection programme for cervical
cancer was established in Puerto Rico in the 1960s.
This covered the metropolitan areas until 1962, and
was later expanded to all health regions of the island.
Cytology smears are offered to women aged
≥15 years and about 150 000 smears are processed
annually. The incidence and mortality from cervical
cancer have declined steadily over the last three
decades (Fig. 1 and Fig. 2). The average, annual age-
standardized incidence dropped from 38 per
100 000 women during 1950–54 to 19.9 per
100 000 women in 1990, and the mortality rate
dropped from 19.1 per 100 000 women to 5.2 per
100 000 women in the same period.

**Sub-Saharan Africa**

There are no organized or opportunistic screening
programmes for cervical cancer in any of the high-
risk sub-Saharan African countries. While data from
Uganda indicate that, at least in some areas of the
country, substantial increases in the incidence of
cervical cancer may have occurred (4), there is no
evidence of an increase in incidence over time in
Zimbabwe (23). Studies in Zimbabwe and South
Africa have assessed the performance characteristics
of potentially alternative screening tests such as visual
inspection with acetic acid (VIA) and HPV testing. A
cross-sectional screening study in Zimbabwe re-
ported that the sensitivity and specificity to detect
high-grade dysplasias and cancer was 77% and 64%,
respectively, for VIA compared to 43% and 91% for
cytology (24). The sensitivity and specificity of HPV
testing using Hybrid Capture II assay (Digene
Corporation, Gaithersburg, USA) were 81% and
62%, respectively (25); the sensitivity and specificity
of HPV testing was, respectively, 91% and 41% for
HIV-infected women and 62% and 75%, respec-
tively, for HIV-negative women (26). It is also
reported that the sensitivity and specificity of VIA
and HPV testing, when used sequentially, was 64%
and 82%, respectively (27).

**South Africa.** The South African Institute of Medical
Research organized the infrastructure for mass
screening of the female population of Soweto
(Project Screen Soweto) so that 90 000 cytology
smears could be tested annually (28). However, the
lack of a planned population education and motiva-
tion programme resulted in poor participation of the
target population in the programme. In a cross-
sectional study that addressed the comparative
performance of cytology, VIA, cervicography, and
HPV testing in South Africa, the sensitivity was
found to be 78%, 67%, 53%, and 73%, respectively;
the specificity was 94%, 83%, 89%, and 86%, respectively (29). In another study in South Africa, HPV testing using self-collected vaginal samples was found to be more sensitive than cytology (66% versus 61%), but less specific (83% versus 88%) (30). In an earlier study in South Africa, the sensitivity of VIA was found to be 65% (31). A recent study of the cost-effectiveness of several cervical cancer screening strategies, based on the South African experience, indicated that strategies using VIA or HPV DNA testing may offer attractive alternatives to cytology-based screening programmes in low-resource settings (32). When all the strategies were analysed on the basis of a single lifetime screening at age 35 years compared with no screening, it was found that HPV testing, followed by treatment of screen-positive women at a second visit, cost US$ 39 per year of life saved (27% reduction in cancer incidence); VIA, coupled with immediate treatment of screen-positive women at the first visit, was the next most cost-effective (26% reduction in cancer incidence) and was cost-saving; cytology, followed by treatment of screen-positive women at a second visit, was the least effective (19% reduction in cancer incidence) at a cost of US$ 81 per year of life saved (32).

Currently, cytology smears are provided on demand in antenatal, postnatal, gynecology, and family planning clinics in South Africa. Work to develop a cervical screening policy for South Africa, based on the models of natural history, has been ongoing for some time. It is proposed to initiate screening at the age of 30 years with three smears being carried out in a woman’s lifetime. However, there has been debate about whether this policy should be implemented and how. A pilot project to set up screening services using the health systems development approach is currently being undertaken by three provincial departments of health (Western Cape, Northern Cape, and Gauteng) in cooperation with nongovernmental organizations. This approach seeks to set up programme components such as reaching the target population, providing a competent screening service, relaying the results, and organizing referral, investigation, treatment and follow-up of screening-positive women. It is expected that these tested methods will be applied in the provinces and then nationally.

A three-arm, prospective randomized intervention trial in South Africa is currently addressing the comparative safety, acceptability and efficacy of screening women with VIA and HPV DNA testing and immediately treating screen-positive women with cryotherapy performed by nurses in a primary health care setting. Outcome measures include reduction of high-grade cervical cancer precursors in treated versus untreated women, followed over a 12-month period.

Other countries. Cross-sectional/randomized screening intervention studies are currently ongoing in several African countries — Burkina Faso, Congo (Brazzaville), Ghana, Guinea (Conakry), Kenya, Mali, Niger, and Nigeria — to address the accuracy of various screening approaches such as cytology, HPV testing, VIA, and visual inspection with Lugol’s iodine (VILI) as well as the detection rates associated with them.

South Asia

India. India accounts for one-fifth of the world burden of cervical cancer. There are no organized or high-level opportunistic screening programmes for cervical cancer in any of the provincial states. Data from population-based cancer registries in different regions indicate a slow, but steady, decline in the incidence of cervical cancer (Fig. 1). However, the rates are still too high, particularly in the rural areas, and the absolute number of cases is on the increase due to population growth. Efforts to improve awareness of the population have resulted in early detection of and improved survival from cervical cancer in a backward rural region in western India (33, 34). Also in two subdistricts of western India where the literacy among women is less than 20% there have been attempts to evaluate the role of improved awareness in the early detection and control of cervical cancer (35). Person-to-person and group health education on cervical cancer were provided to 97 000 women in Madha Tehsil, Solapur district, Maharashtra State, in western India; 79 000 women in Karmala Tehsil served as the control population. This programme was initiated in 1995 and the preliminary results for 1995–99 indicate that, compared with the control area, in the intervention subdistrict a substantially higher proportion of women presented with cervical cancer in earlier stages with significantly reduced case fatality (Table 1).

Visual inspection-based approaches to cervical cancer screening have been extensively investigated in India. The performance characteristics of unaided visual inspection (without acetic acid), also known as “downstaging”, has been addressed in several studies (36). This approach suffers from low sensitivity and specificity to detect cervical neoplasia, particularly the precursor lesions, and is no longer recommended as a screening approach. Currently there are several ongoing, cross-sectional studies being carried out on other screening approaches such as VIA, VIA with magnification (VIAM), and VILI, as well as HPV testing as alternative screening approaches. Results from two reported studies indicate that the sensitivity of VIA to detect high-grade lesions was similar or higher than that of conventional cytology but that its specificity was lower (37, 38).

There are three large, ongoing cluster-randomized intervention trials in India — in Dindigul district (Tamil Nadu), in Mumbai, and in Osmanabad district (Maharashtra) — to evaluate the effectiveness of VIA, in reducing cervical cancer incidence and mortality. The intervention programme in Osmanabad district aims to address the comparative efficacy and cost-effectiveness of three different primary screening approaches in reducing the incidence and mortality: VIA, conventional cervical cytology, and HPV testing. The results of these studies are likely to
provide valuable leads to the development of public health policies to control cervical cancer in developing countries. A recently held national workshop on control of cervical cancer in India reviewed the various methodologies for the early detection of cervical neoplasia and considered both good quality conventional cytology and VIA as suitable tests for early clinical diagnosis (39). In view of the inadequately developed cytology services, VIA was recommended as the immediate option for the introduction of cervical cancer control initiatives as part of the district cancer control programmes in 54 districts in India.

**South-east Asia**

In Singapore, a high level of opportunistic screening for cervical cancer has been operating over the last few years, but has had only minimal impact on the overall incidence and mortality from cervical cancer (3). However, a substantial decline in cervical cancer incidence and mortality has been observed among the Singapore Indian community, with stable trends among the Chinese and Malay communities. Efforts are currently underway to provide an organized screening programme by restructuring the existing opportunistic programme. A test-and-treat approach following VIA is currently being evaluated in Thailand. A cytology-based demonstration programme on screening is currently being implemented by the MOH in Nakornpanam Province in north-east Thailand. The comparative performance of VIA and VILI in detecting cervical neoplasia is being addressed in Vientiane, Lao People's Democratic Republic. Ongoing studies in rural China are addressing the accuracy of cytology and non-cytology-based screening approaches.

**Summary**

Although cytological screening is being carried out in some developing countries/regions, there are no organized programmes and the testing is often of poor quality and performed inadequately and inefficiently among the population. As a result, there has been a very limited impact on the incidence of cervical cancer, despite the large numbers of cytological smears taken in some countries such as Cuba and Mexico. The findings from completed and ongoing research on various approaches to screening (in terms of accuracy and effectiveness) and to treatment (in terms of long-term safety) — such as cryotherapy and loop electrosurgical excision procedures, carried out in field conditions, and test-and-treat approaches — should be taken into account when considering new programmes and when reorganizing existing programmes.

**Effective screening programmes in developing countries**

Efforts to organize effective cervical cancer screening programmes in developing countries will have to find adequate financial resources, develop the infrastructure, train the needed manpower, and elaborate surveillance mechanisms for screening, investigating, treating, and follow-up of the targeted women. Quite often, considerable discussion is focused on which screening test to use — cytology or alternatives to cytology, such as VIA or HPV testing — or which combinations/sequence of screening tests should be used for screening in developing countries. Choosing a suitable screening test is only one aspect of a screening programme. A more fundamental and challenging issue is the organization of the programme in its totality. Whichever screening test is to be used, the challenges in organizing a screening programme are more or less the same. However, screening tests (e.g. cytology, HPV testing) that require additional recalls and revisits for diagnostic evaluation and treatment may pose added logistic difficulties and these may emerge as another barrier for participation in low-resource settings.

The choice of screening test in countries/regions that plan to initiate new programmes should be based on the comparative performance characteristics of cytology and its potential alternatives such as VIA, their relative costs, technical requirements, the level of development of laboratory infrastructure, and the feasibility in a given country/region. Since programmes cannot afford the luxury of frequently repeated testing of women, a highly sensitive test should be provided. If cytology is chosen, considerable attention should be given to obtaining good quality smears, staining, and reporting so that a moderately high sensitivity to detect lesions is ensured. If a potential alternative to cytology, such as VIA, is chosen for screening, considerable attention should be given to the proper monitoring and evaluation of the programme inputs and outcomes before further expansion, since VIA is still an experimental option for cervical cancer screening and it remains to be demonstrated whether VIA-based

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**Table 1. Outcome of information/education on the control of cervical cancer, Solapur District, Maharashtra, India, 1995–99**

<table>
<thead>
<tr>
<th></th>
<th>Intervention area (Madha Tehsil)</th>
<th>Control area (Karmala Tehsil)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total number of women</td>
<td>96 908</td>
<td>76 084</td>
</tr>
<tr>
<td>No. of women–years</td>
<td>352 628</td>
<td>380 805</td>
</tr>
<tr>
<td>No. of incident cervical cancers</td>
<td>80</td>
<td>64</td>
</tr>
<tr>
<td>Stage I and II cancers (%)</td>
<td>65.1</td>
<td>32.8</td>
</tr>
<tr>
<td>Age-standardized incidence (per 100 000)(^a)</td>
<td>26.3</td>
<td>18.7</td>
</tr>
<tr>
<td>No. of deaths from cervical cancer</td>
<td>17</td>
<td>30</td>
</tr>
<tr>
<td>Age-standardized mortality rate (per 100 000)(^b)</td>
<td>5.6</td>
<td>8.6</td>
</tr>
</tbody>
</table>

\(^a\) Incidence rate ratio: 1.41 (95% CI: 1.00–1.98).
\(^b\) Mortality rate ratio: 0.65 (95% CI: 0.36–1.18).
special theme – noncommunicable diseases

screening programmes are associated with a reduction in cervical cancer incidence and mortality. In developing countries, existing ineffective cytology-based programmes should be urgently reorganized and monitored.

Quantitative studies have shown that after two or more negative cytology smears, even screening once every 10 years yields a 64% reduction in the incidence of invasive cervical cancer, assuming 100% compliance (6, 40). Further studies based on this model indicate that once-in-a-lifetime screening may yield around 25–30% reduction in the incidence of cervical cancer (41, 42).

To have an impact on cervical cancer incidence and mortality, efforts must be focused on the following: increasing the awareness of women about cervical cancer and preventive health-seeking behaviour; screening all women aged 35–50 years at least once, before expanding the services and providing repeated screening (e.g. once in every 10 years); providing a screening test with high sensitivity (since women have less frequent opportunities for repeated screening); treating women with high-grade dysplasia and cancer; and monitoring programme inputs and evaluating the outcomes.

Conclusion

Programmes for organized screening of cervical cancer (e.g. in England and Finland) or for opportunistic/spontaneous screening (e.g. in the USA and Canada) involve substantial costs to provide for the associated infrastructure, manpower, consumables, follow-up, and surveillance. In our view, many low-income developing countries, particularly most of those in sub-Saharan Africa, currently have neither the financial and manpower resources nor the capacity in their health services to organize and sustain a screening programme of any sort. Low-income developing countries should consider planned investments in order to improve the capacity of their health services to diagnose and treat cervical cancer precursors and early invasive cancers, before considering even limited screening programmes. VIA may be considered as a suitable early detection test in the context of early clinical diagnosis in low-income countries, particularly in those regions without extensive cytology laboratory facilities.

Those middle-income developing countries with inefficient cytology screening programmes should focus their attention on reorganizing the programme in the light of lessons from their past failures and experiences from elsewhere. Many of these programmes work with the unrealistic notion of offering frequently repeated screening tests (e.g. every year) targeted at women of wide age ranges (e.g. 20–65 years). It would be more realistic and effective to screen high-risk women (e.g. those aged 35–49 years or 30–50 years) only once or twice with a good quality and highly sensitive test, with an emphasis on wide coverage (>80%) of the targeted women. It should also be ensured that women with identified abnormalities attend for diagnosis, management and follow-up. Additional investments should also be made to improve the manpower resources and infrastructure that would sustain the programmes in these countries. Adequate information systems should also be incorporated within the programme for monitoring inputs and outcomes. Middle-income countries without any programmes for cervical cancer screening, but planning to implement such a programme, should consider organizing and sustaining it in a limited geographical region before expanding to cover a wider area. Managerial guidelines are now available to help in planning and implementing appropriate programmes in low-resource settings (13, 43).

Conflicts of interest: none declared.
le dépistage est actuellement inefficace, devront réorganiser leurs programmes à la lumière de l’expérience des autres pays et des leçons de leurs échecs passés. Ceux de ces pays qui envisagent d’organiser un nouveau programme de dépistage devront commencer par une région géographique limitée avant de songer à une quelconque extension. Il est également plus réaliste et plus efficace d’axer le dépistage sur les femmes à haut risque, qui seront soumises une fois ou deux dans leur vie à un test très sensible, en cherchant à obtenir une couverture élevée (>80 %) de la population visée.

Pour organiser un programme de dépistage efficace dans ces pays, il faudra trouver des ressources financières suffisantes, développer les infrastructures, former le personnel nécessaire et élaborer des mécanismes de surveillance pour dépister, examiner, traiter et suivre les femmes appartenant au groupe cible. On tiendra compte des résultats des nombreuses recherches portant sur les diverses approches du dépistage dans les pays en développement ainsi que des directives de gestion existantes lorsqu’on réorganisera des programmes en cours ou que l’on envisagera de nouvelles initiatives en matière de dépistage.

Resumen
Programas eficaces de cribado del cáncer cervicouterino en los países en desarrollo de ingresos bajos y medios
El cáncer cervicouterino representa un importante problema de salud pública entre las mujeres adultas de los países en desarrollo de América del Sur y Centroamérica, el África subsahariana y Asia meridional y sudoriental. Los programas de cribado citológico frecuente, organizados o puntuales, han logrado reducir considerablemente la incidencia de cáncer cervicouterino y la mortalidad asociada en los países desarrollados. En cambio, este tipo de cáncer sigue sin controlarse apenas en los países en desarrollo de alto riesgo, donde las medidas de cribado son ineficaces o inexistentes. El artículo analiza brevemente la experiencia de las iniciativas de cribado e investigación llevadas a cabo actualmente en países en desarrollo.

La infraestructura, los recursos humanos, el material fungible, el seguimiento y la vigilancia que requieren los programas de cribado del cáncer cervicouterino —tanto los organizados como los puntuales— entrañan grandes costos. Debido a lo limitado de sus recursos de atención sanitaria, los países en desarrollo no pueden permitirse el cribado frecuente que durante un amplio intervalo de edades aplican los países desarrollados. Muchos países en desarrollo de bajos ingresos, en particular la mayoría de los países del África subsahariana, no poseen ni los recursos ni la capacidad necesarios para que sus servicios de salud organicen de forma sostenida programa alguno de cribado. Los países en desarrollo de ingresos medios, que aplican hoy medidas de cribado ineficientes, deberían reorganizar sus programas a la luz de las experiencias de otros países y de las lecciones extraídas de sus pasados fracasos. Los países de ingresos medios que decidan organizar un nuevo programa de cribado deberían ensayarlo primero en un área geográfica limitada, antes de estudiar su eventual ampliación. Es más realista y eficaz intentar cribar a las mujeres de alto riesgo una o dos veces a lo largo de su vida mediante una prueba de alta sensibilidad, procurando sobre todo asegurar una amplia cobertura (> 80%) de la población destinataria.

Como parte de las actividades desplegadas para organizar un programa de cribado eficaz en esos países en desarrollo, habrá que hallar recursos financieros suficientes, desarrollar la infraestructura oportuna, capacitar al personal necesario e idear mecanismos de vigilancia para el cribado, investigación, tratamiento y seguimiento de las mujeres destinatarías. A la hora de reorganizar los programas existentes y de planear nuevas iniciativas de cribado, deberán tenerse en cuenta los resultados de las numerosas investigaciones realizadas sobre los diversos enfoques de cribado aplicados en los países en desarrollo, así como las directrices de gestión disponibles.

Referencias


