**Pap and HPV Testing**

**Key Points**

* Cervical cancer screening, which includes the Pap test and HPV testing, is an essential part of a woman's routine health care because it can detect cancer or abnormalities that may lead to cancer of the cervix.
* Current guidelines recommend that women should have a Pap test every 3 years beginning at age 21. These guidelines further recommend that women ages 30 to 65 should have HPV and Pap cotesting every 5 years or a Pap test alone every 3 years. Women with certain risk factors may need to have more frequent screening or to continue screening beyond age 65.
* Women who have received the HPV vaccine still need regular cervical screening.

1. **What causes cervical cancer?**

Nearly all cases of [cervical cancer](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000444973&version=Patient&language=English) are caused by infection with oncogenic, or high-risk, types of human papillomavirus, or HPV. There are about 12 high-risk HPV types. Infections with these sexually transmitted viruses also cause most [anal](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000044185&version=Patient&language=English) cancers; many [vaginal](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000044308&version=Patient&language=English), [vulvar](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000044974&version=Patient&language=English), and [penile](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000257215&version=Patient&language=English) cancers; and some [oropharyngeal cancers](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000446523&version=Patient&language=English).

Although HPV infection is very common, most infections will be suppressed by the [immune system](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046356&version=Patient&language=English) within 1 to 2 years without causing cancer. These transient infections may cause temporary changes in cervical cells. If a cervical infection with a high-risk HPV type persists, the cellular changes can eventually develop into more severe [precancerous](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046220&version=Patient&language=English) [lesions](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046324&version=Patient&language=English). If precancerous lesions are not treated, they can progress to cancer. It can take 10 to 20 years or more for a persistent infection with a high-risk HPV type to develop into cancer.

1. **What is cervical cancer screening?**

Cervical cancer screening is an essential part of a woman’s routine health care. It is a way to detect abnormal cervical cells, including [precancerous](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046220&version=Patient&language=English) cervical [lesions](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046324&version=Patient&language=English), as well as early cervical cancers. Both precancerous lesions and early cervical cancers can be treated very successfully. Routine cervical screening has been shown to greatly reduce both the number of new cervical cancers diagnosed each year and deaths from the disease.

Cervical cancer screening includes two types of screening tests: [cytology](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000335081&version=Patient&language=English)-based screening, known as the Pap test or Pap smear, and HPV testing. The main purpose of screening with the Pap test is to detect abnormal cells that may develop into cancer if left untreated. The Pap test can also find noncancerous conditions, such as infections and [inflammation](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000044042&version=Patient&language=English). It can also find cancer cells. In regularly screened populations, however, the Pap test identifies most abnormal cells before they become cancer.

HPV testing is used to look for the presence of high-risk HPV types in cervical cells. These tests can detect HPV infections that cause cell abnormalities, sometimes even before cell abnormalities are evident. Several different HPV tests have been approved for screening. Most tests detect the [DNA](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000045671&version=Patient&language=English) of high-risk HPV, although one test detects the [RNA](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046568&version=Patient&language=English) of high-risk HPV. Some tests detect any high-risk HPV and do not identify the specific type or types that are present. Other tests specifically detect infection with HPV types 16 and 18, the two types that cause most HPV-associated cancers.

1. **How is cervical cancer screening done?**

Cervical cancer screening can be done in a medical office, a clinic, or a community health center. It is often done during a [pelvic examination](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000322873&version=Patient&language=English).

While a woman lies on an exam table, a health care professional inserts an instrument called a speculum into her vagina to widen it so that the upper portion of the vagina and the [cervix](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046133&version=Patient&language=English) can be seen. This procedure also allows the health care professional to take a sample of cervical cells. The cells are taken with a wooden or plastic scraper and/or a cervical brush and are then prepared for Pap analysis in one of two ways. In a conventional Pap test, the specimen (or smear) is placed on a glass microscope slide and a fixative is added. In an automated liquid-based Pap cytology test, cervical cells collected with a brush or other instrument are placed in a vial of liquid preservative. The slide or vial is then sent to a laboratory for analysis.

In the United States, automated liquid-based Pap cytology testing has largely replaced conventional Pap tests. One advantage of liquid-based testing is that the same cell sample can also be tested for the presence of high-risk types of HPV, a process known as “Pap and HPV cotesting.” In addition, liquid-based cytology appears to reduce the likelihood of an unsatisfactory specimen. However, conventional and liquid-based Pap tests appear to have a similar ability to detect cellular abnormalities.

1. **When should a woman begin cervical cancer screening, and how often should she be screened?**

Women should talk with their doctor about when to start screening and how often to be screened. In March 2012, updated screening guidelines were released by the United States Preventive Services Task Force and jointly by the American Cancer Society, the American Society for Colposcopy and Cervical Pathology, and the American Society for Clinical Pathology. These guidelines recommend that women have their first Pap test at age 21. Although previous guidelines recommended that women have their first Pap test 3 years after they start having sexual intercourse, waiting until age 21 is now recommended because adolescents have a very low risk of cervical cancer and a high likelihood that cervical cell abnormalities will go away on their own.

According to the updated guidelines, women ages 21 through 29 should be screened with a Pap test every 3 years. Women ages 30 through 65 can then be screened every 5 years with Pap and HPV cotesting or every 3 years with a Pap test alone.

The guidelines also note that women with certain [risk factors](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000045873&version=Patient&language=English) may need to have more frequent screening or to continue screening beyond age 65. These risk factors include being infected with the human immunodeficiency virus ([HIV](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000044985&version=Patient&language=English)), being immunosuppressed, having been exposed to [diethylstilbestrol](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046021&version=Patient&language=English) before birth, and having been treated for a precancerous cervical [lesion](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046324&version=Patient&language=English) or cervical cancer.

Women who have had a hysterectomy (surgery to remove the uterus and cervix) do not need to have cervical screening, unless the hysterectomy was done to treat a precancerous cervical lesion or cervical cancer.

1. **What are the benefits of Pap and HPV cotesting?**

For women age 30 and older, Pap and HPV cotesting is less likely to miss an abnormality (i.e., has a lower false-negative rate) than Pap testing alone. Therefore, a woman with a negative HPV test and normal Pap test has very little risk of a serious abnormality developing over the next several years. In fact, researchers have found that, when Pap and HPV cotesting is used, lengthening the screening interval to 5 years still allows abnormalities to be detected in time to treat them while also reducing the detection of HPV infections that would have gone away on their own.

Adding HPV testing to Pap testing may also improve the detection of [glandular](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046386&version=Patient&language=English) cell abnormalities, including [adenocarcinoma](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046216&version=Patient&language=English) of the cervix (cancer of the glandular cells of the cervix). Glandular cells are [mucus](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046274&version=Patient&language=English)-producing cells found in the endocervical canal (the opening in the center of the cervix) or in the lining of the uterus. Glandular cell abnormalities and adenocarcinoma of the cervix are much less common than [squamous cell](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046056&version=Patient&language=English) abnormalities and [squamous cell carcinoma](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046595&version=Patient&language=English). There is some evidence that Pap testing is not as good at detecting adenocarcinoma and glandular cell abnormalities as it is at detecting squamous cell abnormalities and cancers.

1. **Can HPV testing be used alone for cervical cancer screening?**

On April 24, 2014, the Food and Drug Administration (FDA) approved the use of one [HPV DNA test](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000666690&version=Patient&language=English) (cobas HPV test, Roche Molecular Systems, Inc.) as a first-line primary screening test for use alone for women age 25 and older. This test detects each of HPV types 16 and 18 and gives pooled results for 12 additional [high-risk HPV](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000753083&version=Patient&language=English) types.

The new approval was based on long-term findings from the ATHENA trial, a clinical trial that included more than 47,000 women. The results showed that the HPV test used in the study performed better than the Pap test at identifying women at risk of developing severe cervical cell abnormalities.

The greater assurance against future cervical cancer risk with HPV testing has also been demonstrated by a [cohort study](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000285673&version=Patient&language=English) of more than a million women, which found that, after 3 years, women who tested negative on the HPV test had an extremely low risk of developing cervical cancer—about half the already low risk of women who tested negative on the Pap test.

First-line HPV testing has not yet been incorporated into the current professional cervical cancer screening guidelines. Professional societies are developing interim guidance documents, and some medical practices might incorporate primary HPV screening.

1. **How are the results of cervical cancer screening tests reported?**

A doctor may simply describe Pap test results to a patient as “normal” or “abnormal.” Likewise, HPV test results can either be “positive,” meaning that a patient’s cervical cells are infected with high-risk HPV, or “negative,” indicating that high-risk HPV types were not found. A woman may want to ask her doctor for specific information about her Pap and HPV test results and what these results mean.

Most laboratories in the United States use a standard set of terms, called the Bethesda System, to report Pap test results. Under the Bethesda System, samples that have no cell abnormalities are reported as “negative for [intraepithelial](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046336&version=Patient&language=English) [lesion](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046324&version=Patient&language=English) or [malignancy](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000045771&version=Patient&language=English).” A negative Pap test report may also note certain benign (non-neoplastic) findings, such as common infections or [inflammation](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000044042&version=Patient&language=English). Pap test results also indicate whether the specimen was satisfactory or unsatisfactory for examination.

The Bethesda System considers abnormalities of [squamous cells](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046056&version=Patient&language=English) and glandular cells separately. Squamous cell abnormalities are divided into the following categories, ranging from the mildest to the most severe.

**Atypical squamous cells (ASC)** are the most common abnormal finding in Pap tests. The Bethesda System divides this category into two groups, ASC-US and ASC-H.

* + **ASC-US**: atypical squamous cells of undetermined significance. The squamous cells do not appear completely normal, but doctors are uncertain about what the cell changes mean. The changes may be related to an HPV infection, but they can also be caused by other factors.
  + **ASC-H**: atypical squamous cells, cannot exclude a high-grade squamous intraepithelial lesion. The cells do not appear normal, but doctors are uncertain about what the cell changes mean. ASC-H lesions may be at higher risk of being precancerous compared with ASC-US lesions.

**Low-grade squamous intraepithelial lesions (LSILs)** are considered mild abnormalities caused by HPV infection. Low-grade means that there are early changes in the size and shape of cells. Intraepithelial refers to the layer of cells that forms the surface of the cervix. When cells from the abnormal area are removed and examined under a microscope (in a procedure called a biopsy), LSILs are usually found to have mild cell changes that may be classified as mild [dysplasia](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000045675&version=Patient&language=English) or as cervical intraepithelial neoplasia, grade 1 (CIN-1).

**High-grade squamous intraepithelial lesions (HSILs)** are more severe abnormalities that have a higher likelihood of progressing to cancer if left untreated. High-grade means that there are more evident changes in the size and shape of the abnormal (precancerous) cells and that the cells look very different from normal cells. When examined under a microscope, the cells from HSILs are often found to have more extensive changes that may be classified as moderate or severe dysplasia or as CIN-2, CIN-2/3, or CIN-3 (in order of increasing severity). Microscopic examination of HSILs may also reveal [carcinoma in situ](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046488&version=Patient&language=English) (CIS), which is commonly included in the CIN-3 category.

**Squamous cell carcinoma** is cervical cancer. The abnormal squamous cells have invaded more deeply into the cervix or into other tissues or organs. In a well-screened population, such as that in the United States, a finding of cancer during cervical screening is extremely rare.

Glandular cell abnormalities describe abnormal changes that occur in the glandular tissues of the cervix. These abnormalities are divided into the following categories:

**Atypical glandular cells (AGC)**, meaning the glandular cells do not appear normal, but doctors are uncertain about what the cell changes mean.

**Endocervical adenocarcinoma in situ (AIS)**, meaning that severely abnormal cells are found but have not spread beyond the glandular tissue of the cervix.

**Adenocarcinoma** includes not only cancer of the endocervical canal itself but also, in some cases, endometrial, extrauterine, and other cancers.

1. **What follow-up tests are done if cervical cancer screening results are abnormal?**

**For a woman receiving Pap and HPV cotesting:**

If a woman is found to have a **normal Pap test result with a positive HPV test** that detects the group of high-risk HPV types, the doctor will usually have her return in a year for repeat screening to see if the HPV infection persists and whether any cell changes have developed that need further follow-up testing. Alternatively, the woman may have another HPV test that looks specifically for HPV-16 and HPV-18, the two HPV types that cause most cervical cancers.

If either of these two HPV types is present, a woman will usually have follow-up testing with [colposcopy](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000045989&version=Patient&language=English). Colposcopy is the use of an instrument much like a microscope (called a [colposcope](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046465&version=Patient&language=English)) to examine the vagina and the cervix. During a colposcopy, the doctor inserts a [speculum](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046588&version=Patient&language=English) into the vagina to widen it and may apply a dilute vinegar solution to the cervix, which causes abnormal areas to turn white. The doctor then uses the colposcope (which remains outside the body) to observe the cervix. When a doctor performs colposcopy, he or she will usually remove cells or tissues from the abnormal area for examination under a microscope, a procedure called a [biopsy](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000045164&version=Patient&language=English).

If a woman is found to have an **abnormal Pap test result with a negative (normal) HPV test**, the follow-up tests will depend on the Pap test result. If the Pap test result is [ASC-US](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000655176&version=Patient&language=English), the doctor will usually have the woman return in 3 to 5 years for a repeat screen. If the Pap test result is [LSIL](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000410401&version=Patient&language=English), the doctor may recommend colposcopy or might have the woman return in a year for repeat screening.

If a woman is found to have an **abnormal Pap test result with a positive HPV test** that detects any high-risk HPV type, the doctor will usually have the woman receive follow-up testing with colposcopy.

**For a woman receiving Pap testing alone:**

If a woman who is receiving Pap testing alone is found to have an **ASC-US Pap test result**, her doctor may have the sample tested for high-risk HPV types or may repeat the Pap test to determine whether further follow-up is needed. Many times, ASC-US cell changes in the cervix go away without treatment, especially if there is no evidence of infection with high-risk HPV. Doctors may prescribe [estrogen cream](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000753376&version=Patient&language=English) for women with ASC-US who are near or past [menopause](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046296&version=Patient&language=English). Because ASC-US cell changes can be caused by low hormone levels, applying an estrogen cream to the cervix for a few weeks can usually help to clarify their cause.

Follow-up testing for **all other abnormal Pap results** will typically involve a colposcopy.

**For a woman receiving HPV-alone testing:**

If a woman who is having HPV-alone testing **tests positive for HPV types 16 or 18**, she should, according to guidance from the FDA, have a colposcopy. A women who tests negative for types 16 and 18 but is positive for one of the 12 other high-risk HPV types should have a Pap test to determine whether a colposcopy is needed.

1. **How are cervical abnormalities treated?**

If biopsy analysis of cells from the affected area of the cervix shows that the cells have CIN-2 or more severe abnormalities, further treatment is probably needed depending on a woman’s age, pregnancy status, and future fertility concerns. Without treatment, these cells may turn into cancer. Treatment options include the following:

* + LEEP (loop electrosurgical excision procedure), in which an electrical current that is passed through a thin wire loop acts as a knife to remove tissue
  + Cryotherapy, in which abnormal tissue is destroyed by freezing it
  + Laser therapy, the use of a narrow beam of intense light to destroy or remove abnormal cells
  + Conization, the removal of a cone-shaped piece of tissue using a knife, a laser, or the LEEP technique.

The screening guidelines call for women who have been treated for CIN-2 or more severe abnormalities to continue screening for at least 20 years, even if they are over 65.

1. **Do women who have been vaccinated against HPV still need to be screened for cervical cancer?**

Yes. Because current HPV vaccines do not protect against all HPV types that cause cervical cancer, it is important for vaccinated women to continue to undergo routine cervical cancer screening.

1. **What are the limitations of cervical cancer screening?**

Although cervical cancer screening tests are highly effective, they are not completely accurate. Sometimes a patient can be told that she has abnormal cells when the cells are actually normal (a false-positive result), or she can be told that her cells are normal when in fact there is an abnormality that was not detected (a false-negative result).

Cervical cancer screening has another limitation, caused by the nature of HPV infections. Because most HPV infections are transient and produce only temporary changes in cervical cells, overly frequent cervical screening could detect HPV infections or cervical cell changes that would never cause cancer. Treating abnormalities that would have gone away on their own can cause needless [psychological](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000454766&version=Patient&language=English) stress. In addition, follow-up tests and treatments can be uncomfortable, and some treatments that remove cervical tissue, such as [LEEP](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000393533&version=Patient&language=English) and [conization](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046468&version=Patient&language=English), have the potential to weaken the cervix and may affect [fertility](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046399&version=Patient&language=English) or slightly increase the rate of premature delivery, depending on how much tissue is removed.

The screening intervals in the 2012 guidelines are intended to minimize the harms caused by treating abnormalities that would never progress to cancer while also limiting false-negative results that would delay the diagnosis and treatment of a [precancerous](http://www.cancer.gov/Common/PopUps/popDefinition.aspx?id=CDR0000046220&version=Patient&language=English) condition or cancer. With these intervals, if an HPV infection or abnormal cells are missed at one screen, chances are good that abnormal cells will be detected at the next screening exam, when they can still be treated successfully.

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