# Cervical Cancer Prevention: An Update on New Screening and Risk-Based Management Guidelines

Nancy R. Berman MSN, ANP-BC, MSCP, FAANP
Adult Nurse Practitioner/Colposcopist
Certified Menopause Practitioner (NAMS)
Millennium Affiliated Physicians
Division of Michigan Healthcare Professionals
Farmington Hills, Michigan
Clinical Instructor
Department of Obstetrics and Gynecology
Wayne State University School of Medicine
Detroit, Michigan

#### **Disclosures**

Advisory Board: Astellas Speakers Bureau: Astellas

#### Objectives

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- 1. Describe the role of persistent oncogenic HPV in the development of pre-cancer and cancer of the cervix
- List two different uses of HPV testing in cervical cancer screening including co-testing and HPV testing as primary stand-alone screening
- 3. Understand how HPV epidemiology drives risk-based cancer prevention
- 4.Understand why risk-based management represents an improvement in care
- 5.Learn fundamentals of risk-based guidelines for managing patients

#### Goal of Cervical Cancer Screening

 Prevent morbidity and mortality from cervical cancer by:

Identifying and treating high-grade cervical cancer precursors

Avoiding unnecessary and potentially hazardous evaluations and treatment Minimizing costs to healthcare system

Increase benefit and decrease harm!

Saslow D, et al. American Cancer Society, American Society for Colposcopy and Cervical Pathology, and American Society for Clinical Pathology screening guidelines for the prevention and early detection of cervical cancer.

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# Key Facts In the Natural History of HPV

For management and counseling

#### **HPV** and Cervical Cancer

# Virtually all cervical cancers are associated with persistent infection with high-risk HPV types

- Data from a variety of studies have confirmed that certain HPV types are associated with cervical cancer: 16, 18, 31, 33, 35, 39, 45, 51, 52, 56, 58, 59
- Others are probably associated: 26,53, 66, 68,73, 82

Oncogenic HPV is a necessary cause of cervical cancer!

IARC. Monographs on the Evaluation of Carcinogenic Risks to Humans. (in press); Munoz N. Vaccine. 2006.

6

#### Role of Persistent Infection

- Persistent infection with high-risk types of HPV is necessary for the progression of high-grade lesions to invasive cancer
- · Average episode lasts 4-20 months
- <50% of women have same type 1 year later</li>
- Type 16 has a greater risk of persistence

Most HPV infection will go away in a short period of time.

Ho GY, et al. Persistent genital human papillomavirus infection as a risk factor for persistent cervical dysplasia. J Nat Cancer Inst. 1995;87(18):1965-1371. Trotter H. Vaccine. 2006.

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## High Lifetime Risk of HPV Infection

- HPV infections are very common. Nearly everyone will get HPV at some point in their lives.
- More than 42 million Americans are infected with types of HPV that are known to cause disease.
- About 13 million Americans, including teens, become infected each year.

Most everyone who is sexually active will be infected by HPV at some point!

National Center for HIV/AIDS, Viral Hepatitis, STD, and TB Prevention, National Center for Immunization and Respiratory Diseases, National Center for Chronic Disease Prevention and Health Promotion; last reviewed 2/9/24

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**HPV-Associated Disease** 

- · Anogenital cancers
- Cervical
- Anal
- · Vulvar and vaginal
- Penile
- · Other cancers
- · Oral cavity, pharynx, larynx
- Skir
- · Conjunctiva
- · External genital warts
- · Laryngeal papillomatosis

Munoz N. Vaccine. 2006; Lacey CJN. Vaccine.2006.

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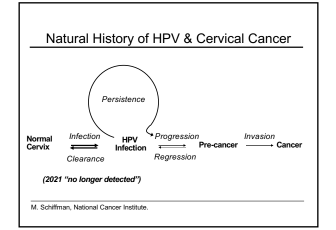
# Risk Factors for *Persistent* HPV Infection and/or Neoplastic Progression

- Smoking
- HPV type
- · Increasing age
- · Lack of condom use
- Immunodeficiency (eg, HIV)
- · Possibly OC use
- · Possibly other STIs, such as chlamydia

Moscicki A-B. Vaccine. 2006; Moscicki A-B. J Infect Dis. 2004; Hogewoning CJ. Int J Cancer, 2003.

Hogewoning CJ. Int J Cancer. 2003.

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## Why Is the Cervix At Risk

**Understanding Transformation Zones** 

14

Moscicki AB. Vaccine, 2006

# Transformation Zones and HPV Infection

- Area where one type of epithelium contacts and gradually replaces another through process of metaplasia
- · Present in cervix, anus, tonsils
- Areas of HPV-related carcinogenesis

Events In Physiologic Metaplasia

EXPOSED COLUMNAR EPITHELIUM

ACID BURN

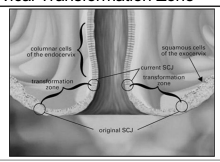
KILLS COLUMNAR RESERVE CELL
EPITHELIUM PROLIFERATION

IMMATURE SQUAMOUS METAPLASIA

EPITHELIAL MATURATION

MATURE METAPLASTIC SQUAMOUS EPITHELIUM

Cervical Transformation Zone



Source: http://www.merckmedicus.com/ppdocs/us/hcp/diseasemodules/hpvd/images/figure25.jpg

New Screening and Risk-Based

Management Guidelines

 Increased knowledge of the natural history of HPV infection has allowed the evolution of screening and management guidelines.

- The role of HPV testing has increased in screening and management .
- There is a paradigm shift from results-based management to risk-based management.
- The management guidelines are available through a phone-based app for purchase or a free web version.

Current Approach to Cervical Cancer Prevention

Requires four separate but linked components:

- HPV vaccination
- Screening
  - Cytology with or without HPV testing Stand alone HPV testing: Primary HPV Screening
- Evaluation of screen-positive women using colposcopy and cervical biopsy
- Treatment of women with biopsy-confirmed highgrade cervical cancer precursors
   Expedited treatment of the highest risk women

rortham EPI, Walf MD, Chuch R. at a Chucia cancer screening for individuals at everage risk 200 guidaire update from the American Cancer Scotely, CAI Cancer J Clin 2007/99(541-548).

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## Change Has Come!

New 2020 American Cancer Society Screening Guidelines Have Changed From 2012!

Current ASCCP, ACP and USPSTF Guidelines for Screening Remain the Same

Evolution in screening: HPV testing --- Cytology--- HPV-· HPV testing detects >90% of precancer More precancer • Pap testing detects <70% after a negative Pap test Screening moving towards HPV testing Less precancer after a negative HPV test Co-testing and HPV primary testing are fairly equivalent HPV self-collection can expand screening FDA approved May 15, 2024 Time since intake testing (months) Dillner, BMJ 2008

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# 2012 ACS/ASCCP/ASCP Cervical Cancer Screening Guidelines



Saslow D, et al. American Cancer Society, American Society for Colposcopy and Cervical Pathology, and American Society for Clinical Pathology screening guidelines for the prevention and early detection of cervical cancer. J Low Genti Tract Dis. 2012;16(3):175-204.

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# Age to Start Cervical Cancer Screening

#### Factors to consider:

- HPV infections are common in young women
- Cervical cancer is rare in adolescents/young women
- Evaluation of minor cytological abnormalities: ls expensive Causes anxiety

Can lead to unnecessary treatments

ACOG Committee on Gynecologic Practice. Obstet Gynecol. 2006.

# Factors Indicating Need for More Frequent Screening

- HIV infection
- Immunosuppression
- DES exposure in utero
- Previous treatment for CIN 2, CIN 3, or cancer

ACOG Practice Bulletin #109. 2009

## Current Cervical Cancer Screening Strategies From the USPSTF and ACS

Patient population	USPSTF (2018)	ACS (2020)
<21 y old	No screening	
21-25 y old	Cytology alone every 3 y	No screening
25-29 y old		Preferred:
30-65 y old	Cytology alone every 3 y     Cotesting b every 5 y     Primary HPV a test every 5 y	Primary HPV a test every 5 y Acceptable: c     Cotesting b every 5 y
		Cytology alone every 3 y
>65 y old	No screening necessary after adequate negative prior screening	
Prior total hysterectomy	No screening necessary in	No screening necessary in
	those without a history of high- grade cervical dysplasia or cervical cancer	those without a history of CIN 2+ or a more severe diagnosis in the past 25 y or cervical cancer ever
Prior HPV vaccination	Follow age-specific recommend:	ations

The ASCCP Cervical Cancer Screening Task Force Endorsement and Opinion on the American Cancer Society Updated Cervical Cancer Screening Guidelines

- The ASCCP recognizes the need to move toward primary HPV-based cervical cancer screening
  - Acknowledges that it will take time to transition clinical and laboratory workflow and operations.
- The ASCCP no longer endorses its 2012 cervical cancer screening guidelines screening that do not include primary HPV
- The combination of abnormal results that occur from either guidance should be managed using the 2019 ASCCP Risk-Based Management Consensus Guidelines.

Marcus, Jenna Z. MD¹, Cason, Pally RN, MS, FNP-BC¹, Downs, Levi S. Jr. MD, MS¹, Einstein, Mark H, MD, MS¹, Flowers, Lisa MD¹ The ASCOP Cervical Cancer Screening Task Force Endorsement and Opinion on the American Cancer Society Updated corrical Cancer Screening Guidelines Journal of Lower General Tract Disease. July 2621 - Volume 27 Sews 3 - p. 187-181 on 1. 10 1/507/LCT 0000000000000014

#### American College of Obstetricians and Gynecologists (ACOG) Practice Advisory 4/21

Replaces Practice Bulletin No. 168, October 2016

- Adoption of the USPSTF guidelines which expands the recommended options for cervical cancer screening in average-risk individuals aged 30 years and older Includes screening every 5 years with primary high-risk human papillomavirus (hrHPV) testing
- Consistent with prior guidance, screening should begin at age 21 years
- Screening recommendations remain unchanged for average-risk individuals aged 21–29 years and those who are older than 65 years
- Management of abnormal cervical cancer screening results should follow current ASCCP guidelines

ACOG (2021). "Practice Advisory: Updated Cervical Cancer Screening Guidelines". from https://www.acog.org/clinical/clinicaguidance/practice-advisory/articles/2021/04/updated-cervical-cancer-screening-guidelines.

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American College of Obstetricians and Gynecologists (ACOG) Practice Advisory 4/21

Screening start age:

- Raising the screening start age to 25 years could: Increase the already high rate of underscreening among individuals aged 25–29 years
   Exacerbate existing health inequities in cervical cancer screening, incidence, morbidity, and mortality
- ACOG, ASCCP, and SGO continue to recommend initiation of cervical cancer screening at age 21 years.

Percent of New Cases by Age Group: Cervical Cancer

Percent of New Cases by Age Group: Cervical Cancer

Cervical cancer is most frequently diagnosed among women aged 35-44.

Median Age At Diagnosis 10 a.1%.

Median Age At Diagnosis 50

SEER 21 2014-2018, All Races, Females

27 28

#### Role of All Professionals....

Advocate for evidenced based guidelines!

# Avoid "OVERPAPULATION" Follow Guidelines!

Neil Lonky ASCCP Biennial Meeting 2008

## **HPV** Testing

## Why Test for HPV?

- Persistent high risk HPV is necessary for the development of cervical cancer
- An obvious corollary is that the absence of HPV means that the risk of cervical cancer is negligible

The negative predictive value for combined HPV Testing and the Pap has been shown to be 99.21% for CIN3.

Sherman ME, et al. J Nat Cancer Inst. 2003;95:46-52.

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# HPV Testing for Screening: Stratifies Risk

- · Allows for less frequent testing
- · Identifies women who need increased surveillance

Wright TC. Obstet Gynecol. 2004. Katki HA et al. Lancet Oncol. 2011.

HPV Detection with FDA-Approved Tests

- Six tests are currently FDA approved and commercially available in the US
- Three are approved for primary, stand-alone screening

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#### FDA Approved HPV Tests

HPV Types Detected	Identifies HPV Type
High and low risk panels (request high risk only)	No
High risk	No (add on test for 16 and 18)
High risk	Yes for 16 and 18
High risk	No (add on test for 16,
	18, and 45)
High risk	Yes for 16,18, 45, 51, 52 Grouped(33, 58, 35, 39, 68
High risk	56, 59, 66)
	Yes 16,18, 45, (31/33/52/58), (35/39/51/56/59/66/68)
	High and low risk panels (request high risk only) High risk High risk High risk

ASCCP. Educate the Educators: HPV and the HPV Vaccines. 2018

...

## **HPV Testing**

FDA approved uses in screening:

• ASC-US Triage

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Preferred age 25-65

Acceptable age 21-24 but does not change management if positive Genotyping is not a factor in management

- Co-testing: Pap and HPV (age 30 and older)
   Genotyping may be used to stratify management
- Primary HPV (Stand alone: age 25 and older)
   FDA approved for only two HPV tests at this time
   Genotyping is reported on all tests

Saslow D, et al. American Cancer Society, American Society for Colposcopy and Cervical Pathology, and American Society for Clinical Pathology screening guidelines for the prevention and early detection of cervical cancer. J Low Genit Tract Dis. 2012;16(3):175-204.

## **HPV Testing**

Other uses of HPV testing:

- · Post abnormal screening and colposcopy follow-up See guidelines1
- Follow-up after cervical treatment

Massad LS, Einstein MH, Huh WK, et al. 2012 updated consensus guidelines for the management of at tests and cancer precursors. Obstetrics and gynecology. 2013;121(4):829-846. 1998;338(7):423-428.

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## 2012 Guidelines: Screening for Women Ages 30-65

- Cytology + hrHPV testing (cotesting) every 5 years is preferred
- Cytology alone every 3 years is acceptable

Saslow D, et al. American Cancer Society, American Society for Colposcopy and Cervical Pathology, and American Society for Clinis screening quidelines for the prevention and early detection of cervical cancer. J Low Genit Tract Dis. 2012;18(3):175-204.

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## ACS/ASCCP/ASCP

"...health care providers can rely on the negative predictive value of the HPV test to assure women who cotest negative that they are at very low risk for CIN3 and cancer for at least 5 years after negative cotesting."

Saslow D, et al. Ca J Clin. 2012.

Women With HPV16 and HPV18 Infections Are More Likely to Develop High-grade Disease

Khan MJ, et al. J Natl Cancer Inst 2005; 97:1072-1079

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Rationale for Cotesting Ages 30-64

• Decreased CIN3 in subsequent screening rounds

· Achieves risk of CIN3 equal to cytology alone

• Enhances detection of adenocarcinoma/AIS · Minimizes the increased number of colposcopies,

• Increased detection of prevalent CIN3

@ 1-3 year intervals

thus it reduces harms.

Co-Testing: Pap and HPV

Women 30 and Older

## Primary HPV Screening

Stand-alone HPV test FDA Approved in 2014 for 25 years and older

Primary HPV:	
Available Testing Platforms	;

Primary HPV test	Year FDA approved	Individual genotypes reported	Pooled genotypes reported	
cobas HPV	2014	16, 18	(31, 33, 35, 39, 45, 51,	
			52, 56, 58, 59, 66, 68)	
Onclarity HPV	2018	16, 18, 31, 45, 51,	(33, 58,	
		52	35, 39, 68	
			56, 59, 66)	
			(31/33/52/58),	
Alinity	2023	16,18, 45	(35/39/51/56/59/66/68)	

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2014 FDA Approval for Primary HPV Testing for Cervical Cancer Screening

#### Rationale

- More sensitive and reproducible than cytology
- · Assesses current and future risk
- · More cost-effective for large-volume screening
- May be more useful in women vaccinated against HPV

Educate the Educator: ASCCP 2016

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## Why HPV Primary Screening?

Co-testing (Pap and HPV testing) is only marginally better than HPV testing alone!

Cox JT, et al. Comparison of cervical cancer screening strategies incorporating different combinations of cytology, HPV testing, and genotyping for HPV 18/18: results from the ATHENA HPV study. Am J Obstet Gynecol. 2013;208(3):184 e181-184 e111.

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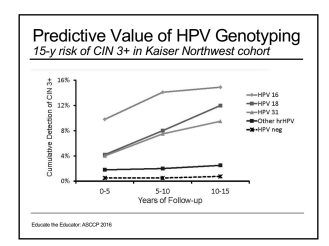
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Importance of Genotyping for HPV 16 &18

- · Over two thirds of cervical cancers in the United States are caused by HPV 16 &18
- Other individual high-risk HPV genotypes are associated with far fewer cancers
- Persistent HPV 16 infection confers a very high risk for CIN 3+, as shown in multiple long-term studies

Wright TC Jr., et al. Evaluation of HPV-16 and HPV-18 genotyping for the triage of women with high-risk HPV+ cytology-negative results. Am J Clin Pathol. 2011;136(4):578-586.

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# ATHENA: Addressing the Need for Advanced HPV Diagnostics

- Prospective, multicenter, US-based study of 47,208 women aged 21 and older
- · Recruited at time of routine screening
- 2.6% had been vaccinated against HPV
- · Screened by liquid based cytology and HPV test

Wright TC, et al. Primary cervical cancer screening with human papillomavirus: end of study results from the ATHENA study using HPV as the first-line screening test. Gynecol Oncol. 2015;138(2):189-197.

Wright TC, Let al. The ATHENA human panillomavirus study: design methods, and baseline results. Am J Obstet Gynecol

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## Absolute Risk of CIN in Cytology: Negative Women ≥ 30 years, Athena Study

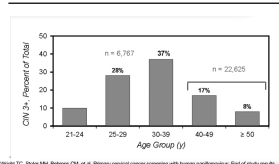
HPV Status	CIN 2+	CIN 3+
High-risk HPV-	0.9%	0.3%
High-risk HPV+ (pooled)	6.3%	4.1%
High-risk HPV 16/18+	11.7%	9.9%
Other 12 High-risk HPV+	4.7%	2.5%

Wright TC, Stoler MH, Behrens CM, et al. Primary cervical cancer screening with human papillomavirus: End of study results from the ATHENA study using HPV as the first-line screening test. Gynecol Oncol. 2015;136(2):189-197. Educate the Educate ASCOP 2016

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## Proportion of CIN3 by Age Group Athena Trial: Why Start Primary Screening at Age 25



Vight TC, Sloler MH, Behrens CM, et al. Primary cervical cancer screening with human papillomavirus: End of study results from the JHTENA study using HPV as the first-line screening test. Gynecol Oncol. 2015;136(2):189-197. Educate the Educator ASCOP 2016.

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2019 ASCCP Risk-Based Management Consensus Guidelines For Abnormal Cervical Cancer Screening Tests

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#### **Original ASCCP Application**



## Risk-Based Management Guidelines

Goals are to increase accuracy and reduce complexity for providers and patients

Development of Guidelines by 19 Participating Organizations

The 2019 guidelines are designed to be enduring, unlike prior versions which required major updates every 5-10 years to adjust for new technologies and emerging evidence.

Perkins RB, Guido RS, Castle PE, et al. 2019 ASCCP Risk-Based Management Guidelines for abnormal cervical cancer screening tests and cancer pre-cursors. J Low Genit Tract Dis. 2020;24(2):102-131

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New Risk-Based Algorithm: Should improve cancer prevention AND decrease unnecessary testing Risk of CIN3+ Provider enters woman's current test results and past 61-100% history 5-60% 1-4% · Risk matrix is used to 0.05% calculate her risk of CIN2/3 0.02% · Computer algorithm <0.01% generates risk score

What Data Were Used/ How Do We Know They Are Representative?

Kaiser Permanente Northern California Data (KPNC)

- Largest/longest real clinical experience with HPVbased screening in the world
- Over 1.5 million women with routine cotesting from 2003-2017
- HPV genotyping for ~19,000 patients
- Provides risk-based evidence for most of the common decision points that occur in screening
- Long length of follow-up allows use of past-history for more personalized management

Cheung LC et al J Low Genit Tract Dis 2020;24(2):90-101

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# Which Risk Factors Influence Pre-Cancer Development?

# HPV Vaccination: Important But NOT Included (yet)!

HPV vaccination prior to age 18 reduces the CIN3+ risk by 50%

#### HOWEVER:

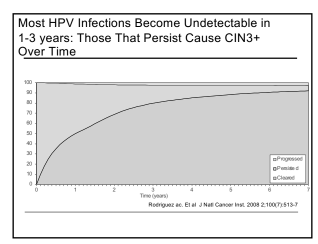
- Current cohort is 21-24 years, a group already conservatively managed.
- 50% age eligible female first dose vaccine population coverage achieved 2015
- Documentation of vaccination and age at which vaccine is necessary to apply this factor correctly—historically guidelines have not included factors clinicians can't document
- Management will likely change as vaccinated cohorts age

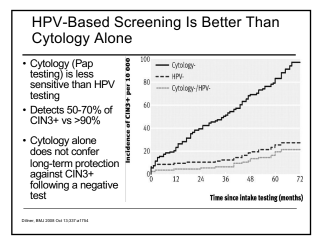
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## Fundamental Concept #1

- The longer an HPV infection has been present, the higher the risk of pre-cancer and cancer
- · Time matters
- Type matters (HPV 16 most dangerous)
- Other patient factors don't matter if you know about HPV
- CLINICAL CORRELATE: Colposcopy is always needed following two consecutive positive HPV tests





# HPV Testing is Necessary for Proper Management

- HPV infections cause nearly all cervical cancers
- · HPV testing is a key component of care
- More precise management is possible when patients are tested

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## New Guidelines Prefer HPV Testing for Follow Up

- Surveillance with cytology alone is acceptable *only* if testing with HPV or cotesting is not feasible.
- Cytology is less sensitive than HPV testing for detection of precancer and is therefore recommended more often.
- Cytology is recommended at 6-month intervals when HPV testing or cotesting is recommended annually.
- Cytology is recommended annually when 3year intervals are recommended for HPV or cotesting.

Perkins RB, Guido RS, Castle PE, et al. 2019 ASCCP Risk-Based Management Guidelines for abnormal cervical cancer screening tests and cancer pre-cursors. J Low Genit Tract Dis. 2020;24(2):102-131

Fundamental Concept #2: Management is Based on Risk, Not Results

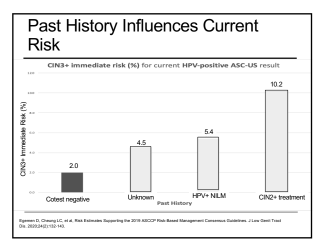
- Recommendations of colposcopy, treatment, or surveillance are based on a patient's risk of CIN3+ determined by a combination of current results and past history (including unknown history).
- The same current test results may yield different management recommendations depending on the history of recent/past test results and other risk factors.

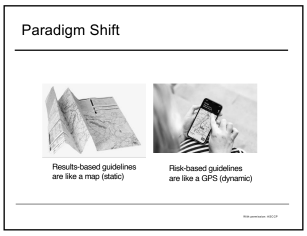
Perkins RB, Guido RS, Castle PE, et al. 2019 ASCCP Risk-Based Management Guidelines for abnormal cervical cancer screening tests and cancer pre-cursors. J Low Genit Tract Dis. 2020;24(2):102-131

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## Risk Thresholds for CIN3 and Management

- Recommendation for colposcopy, treatment, or surveillance is based on a patient's risk of having CIN3+
- This risk is calculated within an algorithm with the patient's current results and any previous results that are available put into an app
- The algorithm is designed to provide the risk-based information with as much or as little previous history as known

Perkins RB, Guido RS, Castle PE, et al. 2019 ASCCP Risk-Based Management Guidelines for abnormal cervical cancer screening tests and cancer pre-cursors. J Low Genit Tract Dis. 2020;24(2):102-131

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# Risk Thresholds for CIN3 and Management

- If the risk for CIN3 is 4% or higher, clinical actions will fall into the categories of colposcopy or expedited treatment
- For patients with a highest risk of 60% or higher, it is preferred to proceed directly to expedited excisional treatment without colposcopy
- Patients with risk between 25% and 59% can choose between expedited treatment or colposcopy

Perkins RB, Guido RS, Castle PE, et al. 2019 ASCCP Risk-Based Management Guidelines for abnormal cervical cancer screening tests and cancer pre-cursors. J Low Genit Tract Dis. 2020;24(2):102-131

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# Risk Thresholds for CIN3 and Management

- In patients with a 4% to 24% risk: colposcopy is preferred
- Patients with a risk below 4% are managed with surveillance: repeat HPV testing or cotesting at 1, 3, or 5 years that is determined by the estimated 5-year CIN3 risk

Perkins RB, Guido RS, Castle PE, et al. 2019 ASCCP Risk-Based Management Guidelines for abnormal cervical cancer screening tests and cancer one-cursors. J Low Genit Tract Dis. 2020;24(2):102-131

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# Patients Stratified Into Risk Levels Look at Immediate CIN3+ Risk for management Look at Freedred Is Immediate CIN3+ risk Expedited Treatment or Colposcopy Acceptable 25-60% immediate CIN3+ risk Colposcopy Acceptable 25-60% immediate CIN3+ risk Colposcopy recommended 4-2-4% immediate CIN3+ risk Colposcopy recommended 4-2-4% immediate CIN3+ risk Return in 1 year 20.15% 5-year CIN3+ risk Return in 5 years <0.15% 5-year CIN3+ risk

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#### 2019 Management Guidelines

Highest Risk Patients Receive Expedited Treatment

- High-grade cytology with HPV16 infections are highest risk
- Excisional treatment for patients at high risk of pre-cancer without requiring confirmatory biopsy

Perkins RB, Guido RS, Castle PE, et al. 2019 ASCCP Risk-Based Management Guidelines for abnormal cervical cancer screening tests and cancer pre-cursors. J Low Genit Tract Dis. 2020;24(2):102-131

## Changes to Follow-up After Treatment of CIN2/3

- HPV-based testing at 6 months, then annually for 3 years
- Continued surveillance with HPV testing or co-testing at 3-year intervals for at least 25 years
- Continued surveillance at 3-year intervals beyond 25 years is acceptable for as long as the patient's life expectancy and ability to be screened are not significantly compromised by serious health issues.

Note: 2012 guidelines recommended return to 5-yr screening intervals and did not specify when screening should cease. New evidence indicates that risk remains elevated for at least 25 yrs, with no evidence that treated patients ever return to risk levels compatible with 5-yr intervals.

ASSUMPTION: Intervals for retesting should reflect underlying risk (equal management for equal risks)

#### The goal was to:

- · Define surveillance intervals
- Define threshold to release patients back to general population screening
- Define risk thresholds for short interval follow up at 1 and 3 years
- Determine which tests to use for surveillance and at what intervals
- HPV alone, HPV/cytology cotesting, cytology (Pap) alone

5-year Return Clinical Action Threshold

#### Guideline:

 When patients have an estimated 5-year CIN3+ risk of <0.15% based on past history and current test results:

Return to routine screening at 5-year intervals using HPV-based testing is recommended.

• Note HPV-based testing is cotesting or primary HPV testing

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#### 3-year Return Clinical Action Threshold

#### Guideline:

 When patients have an estimated 5-year CIN3+ risk ≥0.15% but <0.55% based on past history and current test results:

Repeat testing in 3 years with HPV-based testing is recommended

 Note HPV-based testing is cotesting or primary HPV testing 1-year Return Clinical Action Threshold

#### Guidalina

 When patients have an estimated risk of CIN3+ based on past history and current result that is below the threshold for immediate colposcopy (4.0% immediate risk) and above the 3-year follow-up threshold (≥0.55% at 5 years):

Repeat testing in 1 year with HPV-based testing is recommended

 Note HPV-based testing is cotesting or primary HPV testing

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# Personalized Recommendations Improve Management

- Expedited diagnosis and treatment for high-risk patients
- Fewer invasive procedures on low-risk patients

Patients at High-Risk

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- · Should be referred for expedited treatment
- Specific combinations of test results are so highrisk that patients should proceed directly to a diagnostic excisional procedure (LEEP) HPV 16+ HSIL

HPV-positive HSIL in patients who are underscreened (defined as no screening in more than 5 years)

Perkins RB, Guido RS, Castle PE, et al. 2019 ASCCP risk-based management consensus guidelines for abnormal cervical cancer screening tests and cancer precursors. J Low Genit Tract Dis 2020;24:102–31.

Perkins RB, Guido RS, Castle PE, et al. 2019 ASCCP risk-based management consensus guidelines for abnormal cervic cancer screening tests and cancer precursors. J Low Genit Tract Dis 2020;24:102–31.

# Patients at Medium Risk Should be Referred for Colposcopy

- Patients who are HPV+ twice in a row
- Any HPV16 or HPV18 positive
- Any high-grade Pap result (ASC-H, AGC, HSIL)
   Even if HPV results are negative
- Low-grade Pap results that are HPV positive (ASC-US or LSIL)

Unless preceded by a negative HPV screening test or co-test within 5 years or by a normal colposcopy within 1 year

Perkins RB, Guido RS, Castle PE, et al. 2019 ASCCP risk-based management consensus guidelines for abnormal cervical cancer screening tests and cancer precursors. J Low Genit Tract Dis 2020;24:102–31.

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## The Length of Intensive Surveillance Depends on the Abnormality

- Low-grade abnormalities:
   HPV test or cotest at 1 year
   Extend to 3 year intervals if negative
   Continue for at least 10 years
- CIN2/3 after treatment HPV test or cotest at 6 months, 18 months, 30 months Extend to 3 year intervals if all tests negative Repeat colposcopy for any HPV-positive result

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## Key changes to 2015 Primary HPV Testing Interim Guidance

- HPV 16 or 18 infections have the highest risk for CIN3 and occult cancer, so additional evaluation (e.g., colposcopy with biopsy) is necessary even when cytology results are negative.
- If HPV 16 and 18 testing is positive, and additional laboratory testing of the same sample is not feasible, the patient should proceed directly to colposcopy.

#### Fundamental Concept #3

After an abnormal result, patients enter a surveillance period of close follow up

 All abnormalities require an initial period of intensive surveillance followed by a longer period of surveillance at 3 year intervals

After Initial Surveillance:

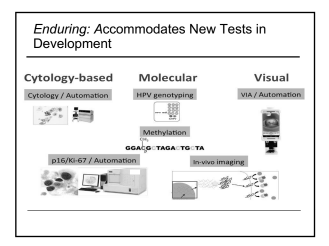
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Screening resumes at 3-year intervals using HPV testing or contesting or annual intervals if using Pap alone

- Surveillance should continue for at least 25 years after treatment for CIN2/2
- · Even if patient undergoes hysterectomy
- Even if patient is over 65
- Screening may continue past age 65 if the patient is in good health

## Enduring: Accommodates New Tests in Development

- Establishment of risk-based thresholds means that new tests can be evaluated against existing thresholds instead of making new algorithms for each new test
- Test characteristics will be objectively compared to existing Clinical Action Thresholds
- Standardized, transparent clinical guidance will logically follow from test characteristics and existing consensus thresholds
- Reduces the need for interim guidance and frequent consensus conferences



#### **Dual Stain**

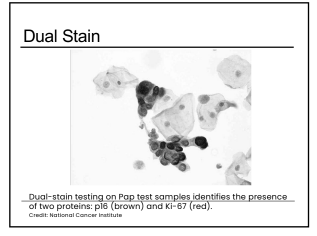
- Identifies abnormal cells through biomarker staining of cervical cells collected during routine screening
- Biomarkers: p16 and Ki-67 when both are positive signal changes at the cellular level that indicate a transforming HPV infection

P16 overexpression is caused by increased E7 oncoprotein activity (correlated with persistent HPV infection)
Ki-67 is a marker of tumor proliferation

- HPV positive women with a negative dual stain could wait 3 years before repeat screening
- HPV positive women with a positive dual stain require colposcopy

"For HPV-Positive Women, Test Can Guide Cervical Cancer Screening Follow-Up", NCI

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Algorithm for Dual Stain Results Patient has HPV+ with no genotyping or with genotyping results negative for 16/18 Patient has any of the following HPV 16 or 18 positive AND ASC-H, HSIL, AGC, or carcinoma No cytology performed or NILM, cytology ASCUS, or LSIL cytology result DS negative Repeat HPV testing Colposcopy recommende DS positive Colposcopy is recommended in 1 year is d (regardless recommended (CIN3+ risk 4.6-11.5%) of DS result) (CIN3+ risk 0.5-0.9%) If results remain DS negative and HPV positive/DS positive, HPV16 or 18 positive, ASC-H, HPV+ with no genotyping or with genotyping results negative for 16/18 HSIL, AGC, or carcinoma and with no cytology performed or cytology → NILM, ASCUS, or LSIL cytology results for 3 consecutive annual Colposcopy recommended results → colposcopy or repeat in 1 ear is acceptable

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## **Extended Genotyping**

- 2 FDA approved tests:
   Onclarity channel configuration
  - HPV 16, 18, 45, 33/58, 31, 52, 35/39/68, 51

Alinity channel configuration

HPV 16,18, 45, 31/33/52/58, 35/39/51/56/59/66/68

- Currently, extended genotyping is being evaluated on whether it will be included into the risk-based management guidelines
  - It is only considering the Onclarity test
    Data for approval was collected with the Onclarity test
    The Alinity test groups the HPV types differently

Self-Collection: HPV Test

## 5/15/24 FDA approved self-collected primary HPV testing in a health-care setting

- Patient uses a collection kit to take a vaginal sample
- · This provides a vaginal HPV test
  - 2 tests were approved
    - Cobas HPV test
  - Onclarity HPV test
- This sample does not allow for cytology or dual stain testing

American Cancer Society Statement: FDA Approval of HPV Self-Collection for Cervical Cancer Screenin

#### Rationale for HPV Self-Collection

- · Increase the number of individuals screened >50% of cancers occur in unscreened/underscreened
- · Reduce barriers to clinician-collected screening Patients may gain access to testing in settings that do not perform speculum exams Discomfort with pelvic exams (mobility, trauma, pain, gender identity)
- · Successful implementation has been seen in programs in The Netherlands, Australia and other sites

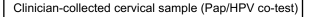
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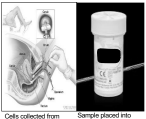
What does self-collection mean for patients, clinicians, and healthcare systems?

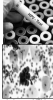
- \*Note, vaginal (self-collected) samples are managed differently than cervical (cliniciancollected) samples
- \*HPV-positive vaginal (self-) samples require speculum exam and Pap collection for evaluation.

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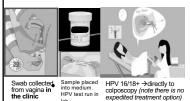


POSITIVE results need colposcopy. Expedited treatment, or 1year follow-up per 2019 ASCCP management quidelines

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Patient-collected vaginal sample (HPV test with second visit for



colposcopy (note there is no expedited treatment option) Other HPV+ → patient returns to office for Pap

Positive Pap →

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#### Implications for patient visits among HPV+

- In a clinician-collected co-testing scenario (our current practice)
- Results returned to clinicians have Pap and HPV results
- Need for colposcopy based on combination of HPV and Pap results
- In a patient-collected vaginal HPV screening scenario, patients with HPV16/18+ proceed directly to colposcopy, all other results require speculum exam for a Pap
- Pap collected at an additional visit with pelvic exam Approximately 60-70% of HPV-positives
- Those with positive results require an additional visit for

Approximately 35-40% of HPV-positives

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Hypothetical population of 100 individuals undergoing self-collection: screening population Visit 1 (or home) 100 Individuals screen with self-collection 90 test negative, N=100 Note may be done in office, laboratory setting +/- primary care visit screening is complete 3 Individuals test 7 Individuals test positive for HPV16 or 18 positive for HR12 Visit 2 N=10 Colposcopy Pap 3 are ASC-US or 4 are NILM 1 year follow-Visit 3 N=3 Colposcopy

## Summary of implications for screening population

- In a self-collection scenario, approximately 90% of patients will test negative, and 10% of patients will test positive and need a follow-up exam
- Among these, 3% go straight to colposcopy, 4% get a Pap test and do not need colposcopy, and 3% need a Pap test and then colposcopy
- Note that self-collection overcomes barriers to screening, but pelvic exams, Pap tests, colposcopy, and treatment are still necessary to prevent cancer

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#### New guidelines coming for self-collection: Focus of guidelines development

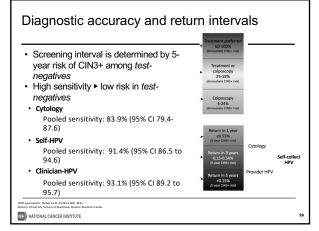
Data analysis:

- Sensitivity of HPV self-collection vs. HPV provider collection vs. cytology: Important for re-testing interval
- Evaluation of overall and type-specific agreement to inform whether extended genotyping recommendations can be applied to self-collected specimens
- Target population for HPV self-collection
- Management of HPV results from self-collected specimens

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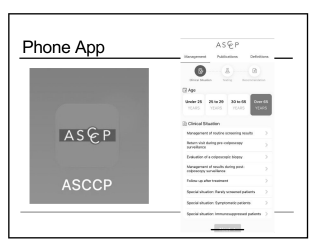
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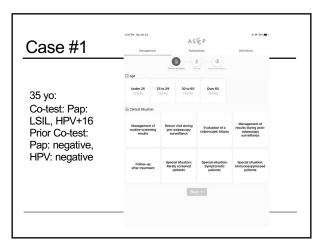
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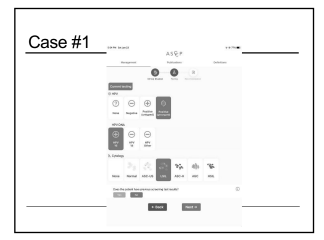


Putting the Risk-Based Guidelines to Use

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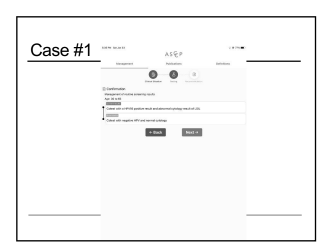






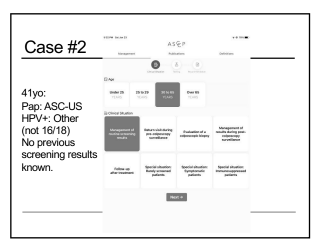


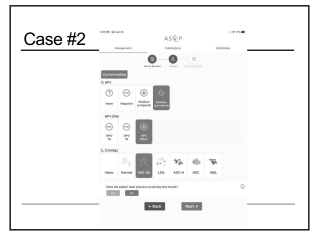
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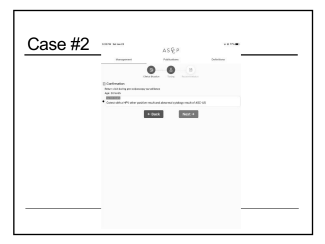


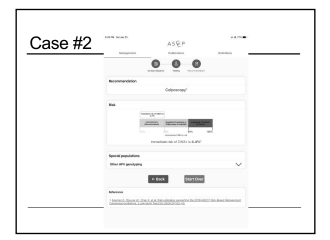


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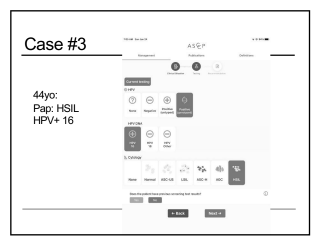


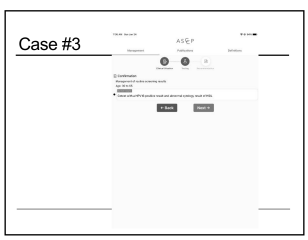






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## Summary

The new guidelines for cervical cancer prevention:

- More benefit with least harm (over screening)
- Identifies low risk women (HPV and Pap negative) and reassures them about safety of longer screening interval
- Identifies truly at-risk women with persistent HPV ... Follow them diligently
- FDA approval of HPV testing as a primary screen, April 2014
- 2019 Risk-based Guidelines: simplify management
- 2019 Risk-based Guidelines: enduring as new tests can be added over time

Never has education of patients and clinicians been more important!

#### Summary

 Majority of cervical cancer in U.S. occurs in women who have not been screened or infrequently screened

Improving access to screening for these women will have a great impact on the prevention of cervical cancer!

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