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Moving forward—the 2019 ASCCP Risk-Based Management Consensus Guidelines for Abnormal Cervical Cancer Screening Tests and Cancer Precursors and beyond: implications and suggestions for laboratories

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*Note: The content of this paper represents the authors' opinions, not necessarily that of the societies they represent.

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Introduction

In 2012, screening and management guidelines for the prevention of cervical cancer adopted the principle of “equal management for equal risk”. Despite the variety of individual test results from different points in the screening and management of an individual, those with similar risks for precancer or cancer can be managed similarly.1 Since 2012, further risk data relevant to cervical cancer have accumulated and show that an individual’s risk of precancer is not dependent solely on their current test results, but may be significantly modified by other factors such as prior screening history, duration and genotype of a human papillomavirus (HPV) infection, and previous treatment for precancer.2-4 Newer approaches to screening and triage have been approved in the United States (eg, primary HPV screening; p16/Ki67 dual staining),7,8 and additional methods and new technologies are either in advanced development or seeking regulatory approval.

The 2012 ASCCP guidelines included 19 management algorithms. The guidelines were complex and did not fully incorporate the increasing understanding of an individual’s risk for cervical precancer. In 2017, the ASCCP, formerly the American Society of Colposcopy and Cervical Pathology, and the National Cancer Institute (NCI) entered into a memorandum of understanding to undertake the development of new risk-based management guidelines that would do just this—incorporate the ability to consider individual-risk by taking into account current test results and past history when managing each patient.9-11 The 2019 ASCCP guidelines highlight that detection and treatment of precancer (high-grade squamous intraepithelial lesion [HSIL]/cervical intraepithelial neoplasia [CIN]3) remains the main aim of cervical cancer prevention in the United States and that the management of patients with abnormal screening results can be optimized by identifying those with the highest risk while minimizing procedures with potential harms for those with low risk.12
A major goal of the 2019 ASCCP guidelines effort was to simplify the management guidelines, thereby clarifying appropriate tests and testing intervals. Risks were estimated by NCI statisticians using large population data sets in the United States. Kaiser Permanente Northern California is the largest, most comprehensive data set in the United States, although for the 2019 guidelines, additional databases were analyzed to ensure that results are applicable to patients of diverse racial, ethnic, and socioeconomic strata. Risk estimates were compared using screening and follow-up data from clinical trials (BD Onclarity registrational trials), the New Mexico State HPV Pap Registry, and the US Centers for Disease Control and Prevention’s National Breast and Cervical Cancer Early Detection Program, a national program that includes many low-income and minority patients.

The risk strata (ranges of risk for CIN 3+) for the guidelines are defined by clinical action thresholds (CATs) that were determined through the consensus process under the principle of “equal management for equal risk”. The immediate and 5-year cumulative risks of CIN 3+ were estimated for each combination of current test results and screening/surveillance history (including unknown history). To determine the next step in management, the estimated risk for a given patient is compared with the proposed CATs (Fig. 1). Management options include return to routine screening, a shortened surveillance interval (1 year or 3 years), colposcopy, or treatment. Providers can use the 2019 ASCCP guidelines to manage their patients by using the tables in Egemen et al or by inputting patient results and history into either a new ASCCP mobile app or Web site designed to facilitate navigation of the tables available at http://www.asccp.org, including a no-cost version. Although there have been requests for integration of the 2019 ASCCP risk-based guidelines with the electronic medical record and laboratory reports, these options are not currently available.

![Decision Diagram](http://www.asccp.org)

**Figure 1** This figure demonstrates how patient risk is evaluated. For a given current results and history combination, the immediate CIN 3+ risk is examined. If this risk is 4% or greater, immediate management via colposcopy or treatment is indicated. If the immediate risk is less than 4%, the 5-year CIN 3+ risk is examined to determine whether patients should return in 1, 3, or 5 years. Adapted with permission from Perkins et al.
The following are examples of potential change to the 2012 “equal management for equal risk” principle of management that will resonate with pathologists as well as providers.

Previously, women with test results of atypical squamous cells of undetermined significance (ASC-US)/HPV-positive were managed with colposcopy—however, based on the 2019 ASCCP guidelines, management may differ depending on prior history/HPV status. The 2019 ASCCP guidelines aim to decrease unnecessary procedures in low-risk women. An example of risk-based management for the same result is illustrated here: An ASC-US/HPV-positive result will have an immediate risk of CIN 3+ of 4.4% (exceeding the 4% colposcopy threshold) when the prior history is unknown. However, if the prior result was a negative HPV primary screen or cotest, the immediate CIN 3+ risk is 2% if, resulting in a 1-year HPV-based follow-up (Table 1A, Egemen D, et al). If the ASC-US/HPV-positive was followed by colposcopy with biopsy showing CIN1 the prior year, the negative colposcopy reduces the risk of CIN 3+ within the next 2 years, and the immediate risk of CIN 3+ would be 3.1%, also resulting in a 1-year HPV-based follow-up (Table 1A, Egemen D, et al).

The New Technology Committee will continue to assess and implement updates as new tests become approved and available, providing an enduring platform for new developments to be incorporated and the ability for the 2019 ASCCP guidelines to be updated on a more regular basis, without a large consensus conference effort. Additional risk modifiers, such as HPV vaccination status, will also be added, as vaccinated cohorts in the United States enter screening age, and enough data accumulates. A deliberative method of stakeholder engagement was utilized in the development of these consensus guidelines, with the inclusion of representatives from 19 stakeholder organizations in the guideline working groups. Input from a number of surveys of providers and patients, and feedback from an open public comment period on draft guidelines, were considered before finalizing the 2019 ASCCP guidelines. Pathologists were well represented by major professional organizations, with 11 pathologists distributed among the working groups and 2 on the steering committee. The 2019 ASCCP Risk-Based Consensus Management Guidelines rolled out on April 2, 2020 during the virtual ASCCP annual meeting.

As laboratorians, how can we support cervical cancer prevention in view of the new screening and management guidelines? Herein we summarize the changes in the 2019 ASCCP guidelines that are relevant to laboratories, pathologists, and cytotechnologists. Prior relevant screening and reporting recommendations that have not been widely and/or consistently adopted by laboratories are also discussed. We also offer considerations for modification of laboratory practices as we transition to these updated practice guidelines.

Human papillomavirus (HPV) testing

HPV has long been established as a necessary cause of cervical cancer. Over the past 3 decades, advances in our understanding of the pathobiology of HPV have clarified that it involves the anogenital tract epithelium in 2 distinct ways: HPV infection and true precancer. In the context of HPV testing for cervical cancer prevention, our goal is to detect those infections that may progress to precancer or cancer; hence, only testing platforms detecting high-risk or oncogenic types of HPV should be used. Low-risk types of HPV infection can cause external genital warts and are rarely, if ever, linked to cervical cancer or high-grade precursor lesions. Testing for low-risk HPV infections in the cervix does not provide information with regard to a patient’s risk status and/or impact patient management. In fact, identification of low-risk HPV infection may lead to additional unnecessary testing and procedures, which can result in patient harm. Thus, we recommend against testing for low-risk HPV types.

HPV testing was introduced into the repertoire for cervical cancer screening and management in the United States in the mid 1990s. The indications for HPV testing vary based on the specific clinical situation for each patient and include screening, triage, and surveillance. Use of high-risk HPV testing was endorsed in screening and management guidelines, initially as a triage test for the cytologic result of ASC-US (2001). Regulatory approvals for cotesting (2003), for post-colposcopic/post-treatment follow-up and risk stratification using partial genotyping (2006), and as a stand-alone primary screening option (2014) followed.

In 2018, the US Preventive Services Task Force endorsed primary HPV screening for women aged 30 to 65 years. The updated American Cancer Society guidelines, due to be released in summer 2020, are also likely to endorse primary HPV screening in the United States. Although the United States has not widely adopted primary HPV testing as a screening modality, this transition is currently underway. These ongoing updates to cervical cancer screening guidelines, both in the United States and internationally, have been based on the higher sensitivity of HPV-based screening, modeling studies, and data from a number of trials, both in the United States and internationally. Comparison of screening methods is not the primary subject of this publication; however, additional communications, including from the Cytopathology Education and Technology Consortium, discuss concerns regarding the approval and implementation of primary HPV screening as the preferred screening strategy, without the option of cotesting, in the current US opportunistic screening program.

There are currently 5 HPV tests approved by the US Food and Drug Administration (FDA) for cytology and HPV cotesting and reflex indications: Qiagen Hybrid Capture (Gaithersburg, MD), Hologic Cervista and Hologic Aptima (Marlborough, MA), Roche cobas (Indianapolis,
<table>
<thead>
<tr>
<th>Test</th>
<th>Hybrid capture II</th>
<th>Cervista</th>
<th>Cobas</th>
<th>Aptima</th>
<th>BD Onclarity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manufacturer</td>
<td>Qiagen</td>
<td>Hologic</td>
<td>Roche</td>
<td>Gen Probe (Hologic)</td>
<td>Becton Dickinson</td>
</tr>
<tr>
<td>Year FDA approved for reflex HPV testing and HPV/Papanicolaou cotesting</td>
<td>2001</td>
<td>2009</td>
<td>2011</td>
<td>2011</td>
<td>2018</td>
</tr>
<tr>
<td>Year approved for primary screening</td>
<td>N/A</td>
<td>N/A</td>
<td>2014 (ThinPrep only)</td>
<td>N/A</td>
<td>2018 (SurePath only)</td>
</tr>
<tr>
<td>Method</td>
<td>DNA (non-PCR based)</td>
<td>DNA (non-PCR based)</td>
<td>DNA (PCR based); Target amplification: L1 gene target</td>
<td>mRNA (PCR based); Target amplification: E6/E7 gene target</td>
<td>DNA (PCR based); Target amplification: E6/E7 gene target</td>
</tr>
<tr>
<td>Genotypes detected</td>
<td>16, 18, 31, 33, 35, 39, 45, 51, 52, 56, 58, 59, 68</td>
<td>16, 18, 31, 33, 35, 39, 45, 51, 52, 56, 58, 59, 66, and 68</td>
<td>16, 18, 31, 33, 35, 39, 45, 51, 52, 56, 58, 59, 66, and 68 with genotyping of 16 and 18</td>
<td>16, 18, 31, 33, 35, 39, 45, 51, 52, 56, 58, 59, 66, and 68; genotyping as separate test (16, 18/45)</td>
<td>16, 18, 31, 33, 35, 39, 45, 51, 52, 56, 58, 59, 66, 68; simultaneous, discrete identification of 16, 18, and 45</td>
</tr>
<tr>
<td>Clinical trial</td>
<td>ASC-US/LSIL Triage Study (ALT5), 2006 CAP</td>
<td>Cervista HPV HR</td>
<td>ATHENA</td>
<td>CLEAR trial</td>
<td>Onclarity trial (baseline phase)</td>
</tr>
<tr>
<td>Clinical validation</td>
<td>Extensive</td>
<td>Limited</td>
<td>Limited</td>
<td>Limited</td>
<td>Limited</td>
</tr>
<tr>
<td>Sensitivity for CIN 2/3</td>
<td>63.6%-100%</td>
<td>92.8%-100%</td>
<td>71.1%-99%</td>
<td>55.3%-100%</td>
<td>85.7%-100%</td>
</tr>
<tr>
<td>Specificity for CIN 2/3</td>
<td>6.2%-98.4%</td>
<td>—</td>
<td>24%-86.2%</td>
<td>28.8%-99.2%</td>
<td>17%-98.8%</td>
</tr>
<tr>
<td>Built-in internal control</td>
<td>No</td>
<td>Yes (HIST2H2BE)</td>
<td>Yes (ß-globin)</td>
<td>Yes (ß-globin)</td>
<td>Yes (ß-globin)</td>
</tr>
</tbody>
</table>

Abbreviations: N/A, not applicable; PCR, polymerase chain reaction.
Adapted with permission from Salazar et al.24
IN), and Becton Dickinson Onclarity (Franklin Lakes, NJ). All of these tests, except Aptima, utilize DNA hybridization and amplification or signal amplification. The Aptima test uses RNA amplification and some studies suggest that it is slightly more specific for high-grade precancers and cancers.23 All, except Qiagen’s HC2, have an internal control; however, none of the internal controls specifically identify epithelial cells. Only 2 of these platforms are approved for primary screening: cobas and Onclarity. The 2015 ASCCP interim and new (2019) guidelines, along with the Cytopathology Education and Technology Consortium, recommend that only platforms approved for primary screening be used when HPV alone is performed as a follow-up test strategy for women with a history of abnormal screening tests.15,22 Table 1 summarizes the characteristics of the FDA-approved HPV testing platforms currently available in the United States.24

In the 2019 ASCCP risk-based management consensus guidelines, HPV-based testing is the basis for risk estimation and laboratories will likely see a larger volume of HPV testing requests.13,14 The term HPV-based testing is used in the 2019 ASCCP guidelines to refer to use of either primary HPV testing alone or HPV testing in conjunction with cervical cytology (cotesting).15 Previously, HPV testing had mostly been used for cotesting or as a reflex test for an ASC-US cytology.

The 2019 ASCCP management guidelines15 specifically state:

- Human papillomavirus assays that are Food and Drug Administration (FDA)-approved for screening should be used for management according to their regulatory approval in the United States. (Note: all HPV testing in this document refers to testing for high-risk HPV types only).
- For all management indications, HPV mRNA and HPV DNA tests without FDA approval for primary screening alone should only be used as a cotest with cytology, unless sufficient, rigorous data are available to support use of these particular tests in management.
- 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 pre-cancer and is therefore recommended more often.

**HPV genotyping**

The 2019 ASCCP guidelines incorporate partial genotyping (HPV 16/18) information, if available, in the risk assessment profile for patients who test positive for high-risk HPV types. This guidelines consensus effort acknowledged that not all laboratories may currently have genotyping capability; however, when available, this additional information further stratifies risk.15 Large data sets analyzed as part of the guidelines review process show that the identification of HPV 16 clearly mandates consideration in clinical management of new abnormal screening results. HPV 18 positivity must be considered as a special situation because of an established disproportionate risk of invasive cancer.25

When HPV-based screening is performed, the 2019 ASCCP guidelines recommend that:

- All positive primary HPV screening tests, regardless of genotype, should have additional reflex triage testing performed from the same laboratory specimen (e.g., reflex cytology). This is a change from the 2015 ASCCP interim guidance for management of primary HPV screening results.22
- Additional testing from the same laboratory specimen is recommended because the findings may inform colposcopy practice. For example, the CAT for expedited treatment is met for patients who test HPV-16 positive with HSIL cytology; this approach bypasses the intermediate step of colposcopic biopsy confirmation of high-grade disease.
- If HPV 16 or 18 testing is positive, and additional laboratory testing of the same sample is not feasible, the patient should proceed directly to colposcopy.
- HPV 16 or 18 infections have the highest risk for CIN 3 and occult cancer, so additional evaluation (e.g., colposcopy with biopsy) is necessary even when cytology results are negative or unavailable.

Laboratory requisitions and electronic medical records/laboratory information system ordering options used at this time are likely to have been developed predominantly for HPV cotesting and reflex testing. Cytopathology professionals and laboratories should be prepared to facilitate testing strategies recommended in the 2019 ASCCP guidelines by assessing the following:

(a) Ability to identify and confirm primary versus cotest HPV testing requests in a clinical context.
(b) Ability to offer FDA-approved primary HPV screening and genotyping platforms.
(c) Testing options/order sets and workflow for HPV based screening/management options within the cytopathology laboratory and other locations (e.g., molecular diagnostics/referred testing).
(d) Integrated reporting of results (preferred), when both HPV and cytology are utilized, whether as cytology/reflex HPV, cotesting, or primary HPV/reflex cytology and irrespective of the test indication (screening, triage or management).
(e) Electronic medical record clinical decision support implementation, especially passive clinical decision support, to guide age-compliant screening and triage options and support clinical providers.

Clinical providers are equally obligated to provide accurate and sufficient patient history on the requisition form to allow the laboratory to make certain that HPV test orders are consistent with the 2019 ASCCP guidelines. Providers should also ascertain that the HPV testing platform being utilized by their laboratory is FDA-approved for the specific test indication. Cytopathology professionals and laboratories should discuss
HPV testing platforms with clinicians and monitor compliance with age-specific screening and risk-based management guidelines and provide feedback to their clinician groups.

**Reporting and management of anatomic pathology results**

**Cytopathology**

**Atypical glandular cells (AGC) and adenocarcinoma in situ (AIS)**

The Bethesda System (TBS) 2001 update replaced the term “atypical glandular cells of undetermined significance” (AGUS) with the term atypical glandular cells (AGC), and recommended a 2-tiered subcategorization: first, as to cell type/origin, if possible (ie, endocervical versus endometrial, or glandular when uncertain), and second, as favor neoplasia or not otherwise specified (NOS) for AGC and atypical endocervical cells (AEC). The rubric of AGC also includes atypical endometrial cells. Although cervical cytology is not a screening test for endometrial pathology, lesional cells can sometimes be seen. Atypical endometrial cells are not further subcategorized in terms of risk of neoplasia. TBS 2001 also introduced a specific cytologic interpretation of adenocarcinoma in situ (AIS). There were no changes made to the reporting categories for glandular epithelial cell abnormalities in TBS 2014.

The overall category of AGC, albeit rare, may correspond histologically to several different entities ranging from benign (eg, polyps, tubal metaplasia) to intraepithelial lesions and invasive cancer, and include both glandular and squamous lesions (eg, HSIL involving endocervical glands, AIS, invasive adenocarcinoma of the Mullerian tract or other sites). AGC is a much higher-risk cytologic interpretation than ASC-US. Even with established TBS diagnostic criteria, cytologic glandular abnormalities remain poorly reproducible and diagnostically challenging; this results in both under- and over-interpretation by cytotechnologists and cytopathologists. Some laboratories target AGC cases for consensus review by additional pathologists prior to issuing a final report.

The 2019 ASCCP guidelines for glandular cytologic abnormalities remain similar to the 2012 guidelines. Clinicians and cytologists often incorrectly think of cytologic AGC as a group with the same management. Triage of any category of AGC by HPV testing continues to be unacceptable in the 2019 ASCCP guidelines. All cytologic glandular abnormalities will lead to initial colposcopic evaluation with endocervical sampling except when atypical endometrial cells are specified. In that case, the preferred initial evaluation is endometrial and endocervical sampling. Additional endometrial sampling is also recommended when the patient is 35 years of age or older, symptomatic, or at high risk for endometrial neoplasia based on risk factors. The 2019 ASCCP guidelines emphasize that although endometrial cancer is rare in premenopausal patients without risk factors, the prevalence of premenopausal endometrial cancer is increasing, underscoring the importance of endometrial sampling when indicated. Subsequent management for cytologic AEC/AGC, favor neoplasia and AIS is a diagnostic excision procedure, even if a high-grade abnormality is not confirmed on colposcopic biopsy with endocervical sampling, and even if HPV testing is negative. As endocervical lesions are difficult to visualize clinically, colposcopists rely on pathologists to provide appropriate categorization of cell origin and level of suspicion for neoplasia so that appropriate management is possible.

**Atypical squamous cells, cannot exclude HSIL (ASC-H) and combinations of cytology results**

According to the Kaiser Permanente Northern California data, HPV-negative atypical squamous cells, cannot exclude HSIL (ASC-H) and HPV-positive ASC-H had very different CIN 3+ rates, but similar cancer rates. HPV-positive ASC-H had an immediate CIN 3+ risk of 26% and a cancer risk of 0.92%, whereas HPV-negative ASC-H had an immediate CIN 3+ risk of 3.4%, but an immediate cancer risk of 0.69%. Because the immediate cancer risk for ASC-H is disproportionately high compared with the CIN 3+ risk, the 2019 guidelines carried forward the 2012 recommendations of colposcopy for all patients with ASC-H, regardless of HPV test results.

HSIL found in the background of atrophy is often difficult to appreciate because of the lack of maturation of squamous cells and the similarity between small atrophic cells and the dysplastic cells. Additionally, as illustrated in the 2014 Bethesda Atlas, isolated highly atypical squamous cells, with very large nuclei, a characteristic smudgy or degenerative chromatin pattern, and a very high nucleus to cytoplasmic ratio can be occasionally identified in deeply atrophic specimens. Such cells can elicit morphologic concern for HSIL, but are observed, often in older patients with few or no risk factors. A conservative approach, such as interpretation as ASC-US with follow-up/reflex HPV testing, rather than ASC-H, may be more appropriate in such cases. In cases of atrophy with abnormal cells meeting criteria for HSIL, however, an interpretation of HSIL should be made.

When reporting squamous intraepithelial lesions (SIL) according to Bethesda terminology, it is important to recognize that the concurrent presence of LSIL cells is not necessary to make an interpretation of HSIL and the presence of even a small population of definitive HSIL cells in the background of a predominance of LSIL cells should result in an interpretation of HSIL. In rare cases, cytology specimens may exhibit squamous cells with features that lie between low- and high-grade SIL. These include cases that exhibit predominantly LSIL and a few cells suggestive, but not diagnostic, of HSIL (ASC-H) or cases that display...
keratinized cells with dense eosinophilic cytoplasm that give an impression of higher nucleus to cytoplasmic ratio than in classic LSIL, but without specific features of classic HSIL.  

Per TBS 2014, such cases should be reported using well-established TBS categories, specifically “ASC-H in a background of LSIL”, and not in other terms such as “LSIL, cannot exclude a high-grade lesion”.  

Non-TBS reporting terms do not provide clear management direction to the clinical provider, since they are not linked to the ASCCP management guidelines as are the TBS categories. Management under the 2019 ASCCP guidelines is based on the highest-risk cytologic interpretation, for example, ASC-H, when both ASC-H and LSIL are reported on the same sample. In other rare combinations of cytologic results such as HSIL and AGC, or ASC-US and AGC, management is also determined by the highest risk abnormality.

### Histopathology

#### Reporting of HPV-Associated squamous intraepithelial lesions

Both the Lower Anogenital Squamous Terminology (LAST) Project and the World Health Organization recommend the use of a 2-tiered nomenclature for HPV-associated squamous cervical lesions: LSIL, the morphologic representation of HPV infection, and HSIL, the histopathological correlate of precancer. Rather than the spectrum of disease that is implied by the 3-tiered CIN classification system, the 2-tiered system of LSIL/HSIL better reflects the known biology of HPV disease. Both LAST and WHO also permit the CIN grade to be designated in parenthesis after the SIL nomenclature. Adoption of this terminology by pathologists has not been universal.

The 2019 ASCCP guidelines include the following specific statement regarding the use of p16 immunohistochemistry (IHC) and a 2-tiered nomenclature for reporting histopathology of HPV-associated squamous lesions of the lower anogenital tract.

- It is important to use p16 immunohistochemical staining according to the guidance provided by the College of American Pathologists (CAP)-ASCCP Lower Anogenital Squamous Terminology (LAST) Project. p16 immunohistochemistry should be used for specific indications as recommended by the LAST guidelines when interpreting the hematoxylin and eosin (H&E) slide. A positive p16 immunostain supports the diagnosis of histologic HSIL if the morphological assessment of H&E slides is consistent with CIN 2 or CIN 3. There is a risk of overcalling cervical histology findings when p16 is used incorrectly. Most importantly, a morphologic CIN1 on H&E should not be upgraded to histologic HSIL (CIN 2), even if p16 positive.

- For epidemiologic and clinical management purposes, it is strongly recommended to qualify a histologic HSIL result by CIN 2 or CIN 3, according to the options given by the LAST guidelines (eg, histologic HSIL [CIN 2]).

### Rationale

This CIN qualification can have clinical importance (eg, to identify cases of CIN 2 in patients for whom conservative management is an acceptable option). It is also important for post-vaccine surveillance studies and quality control assessments of cervical precancer that have historically relied on CIN 2 and CIN 3 endpoints. Furthermore, it is important for future research efforts to distinguish diagnoses of histologic HSIL (CIN 2) from HSIL (CIN 3) so that diagnostic categories are compatible with the histologic endpoints used for current guidelines.

### Note

Strong and diffuse block staining for p16, as defined by LAST is “continuous strong nuclear or nuclear plus cytoplasmic staining of the basal cell layer with extension upward involving at least one third of the epithelial thickness”. The latter height restriction is somewhat arbitrary but adds specificity. Full-thickness staining or extension into the upper third or upper half is specifically not required to call a p16 immunostain positive. Focal or patchy staining is nonspecific and can be seen with reactive squamous metaplasia, as well as low-grade disease (LSIL, CIN 1). All other staining patterns, described as cytoplasmic only, wispy, scattered, single cells, and others, are defined as negative.

It is important for pathologists to remember that p16 IHC is not a highly specific test and thus to use p16 only as suggested by the LAST recommendations to avoid over-usage. As with other immunostains, correct interpretation of p16 IHC is also needed to avoid over interpretation of positivity. The LAST recommendations are undergoing an update, starting in 2020, and additional biomarkers could possibly further refine morphologic interpretations.

The 2019 ASCCP guidelines recommend treatment of histologic HSIL in most circumstances in non-pregnant women. Treatment is specifically recommended for HSIL (CIN 3), for any HSIL if the proximal extent of the lesion is in the endocervical canal or the entire squamocolumnar junction cannot be visualized, or if endocervical sampling shows HSIL or ungraded CIN. Treatment is also recommended for HSIL not further qualified (ie, CIN 2/3). Observation is preferred only when HSIL is specifically qualified as CIN 2 and only in specific situations: for women <25 years or for women whose concerns about pregnancy-related treatment complications outweigh concerns about cancer. Thus, a pathology report of cervical HSIL, regardless of CIN grade or unqualified, serves as the foundation for treatment decisions in most situations.

Given that background, how can pathologists give the most reliable histopathology diagnoses for cervical biopsies to their clinical colleagues to facilitate sound patient management recommendations? Any morphologic interpretation is subject to interobserver variation. Pathologists give more consistent diagnoses on cervical biopsies with the judicious
use of p16 IHC. Improved interobserver agreement for the diagnosis of CIN 2+ with the conjunctive use of hematoxylin and eosin (H&E) morphology and p16 IHC compared with H&E morphology alone has been shown in several studies,\textsuperscript{35-38} including in a systematic review and meta-analysis.\textsuperscript{39} Because histologic HSIL is the usual trigger for treatment, the use of p16 per the LAST and WHO guidelines provides our clinical colleagues more reproducible diagnoses upon which to base management recommendations.

The LAST Project recommends that p16 IHC, when positive, should be used to support the diagnostic interpretation of HSIL in times of diagnostic uncertainty (such as with atypical squamous metaplasia on H&E) and to support a diagnosis of HSIL (CIN 2). CIN 2 has historically been the threshold for treatment in most circumstances. However, among pathologists, CIN 2 is the least reproducible of the CIN categories.\textsuperscript{30,44} If a diagnosis of CIN 2 is entertained based on solid morphologic criteria, a p16 should be performed to support a high-grade lesion. If negative, the diagnosis should be downgraded to LSIL or a benign process depending on the differential diagnosis on H&E. Judicious use of p16, in conjunction with H&E morphology is needed, however. When the morphologic diagnosis is LSIL (CIN 1), a p16 should not be performed. Morphologic LSIL (CIN 1) is frequently p16-positive; thus if p16 is positive, LSIL should not be upgraded to HSIL. There is insufficient data to indicate that the risk profile of p16-positive LSIL meets the CAT for treatment. Given diagnostic uncertainty encountered in tissue diagnoses of challenging histomorphology and the inherent interobserver variability of morphologic diagnosis, p16 IHC, when positive, is a useful tool to support a diagnosis of precancer.

More high-grade lesions may be identified as clinicians adopt the ASCCP colposcopy standards published in 2017.\textsuperscript{42} These standards recommend that colposcopists take multiple biopsies of all areas with acetowhitening, metaplasia, or higher colposcopic abnormalities. As a result, many colposcopic procedures will have at least 2 and up to 4 targeted biopsies from distinct lesions. This recommendation should help reduce sampling issues related to colposcopic-directed biopsies and interobserver variability of colposcopic impression of lesion grade, but it may also detect more small lesions that are morphologically high-grade.

An important and potentially overlooked issue is that of the size of the cervical HSIL. Lesion size is not a component of the treatment algorithms; treatment recommendations are based on the histopathology diagnosis of a potential precancer regardless of lesion size. In the only known and now infamous “Unfortunate Experiment”, an observational study of untreated CIN 3, the long-term risk of developing invasive cancer was approximately 30% over a 30-year period.\textsuperscript{43} As Schiffman and Rodriguez accurately point out in an accompanying commentary, these were large, prevalent lesions in women with a median age of 38 years.\textsuperscript{44} The natural history of early, small lesions, potentially detected only because of increased sampling and screening efforts, remains unknown.

Another issue is the inability to accurately grade a fragmented squamous intraepithelial lesion (SIL), which is often reported as SIL, ungraded. Newer cervical biopsy collection devices, such as the SoftBiopsy, (Histologics Anaheim, CA; https://histologics.com/softbiopsy.html) and endocervical curettage specimens “denude” the cervical epithelium for tissue examination rather than provide a full-thickness tissue biopsy. This can result in problems with tissue orientation that make the lesion difficult to grade. These cases are more likely to result in a diagnosis of “SIL, ungraded”. A recent study by Lee et al\textsuperscript{45} found that histologic SIL, ungraded, although representing only 1.9% of their total cases, resulted in positive results (HSIL, AIS, or carcinoma) in 41% of patients with histologic follow-up. They found that a combination of p16 and ki-67 helped to correctly classify these lesions. Many clinical care providers do not understand that ungraded SIL represents a high-risk category for precancer and patients may not undergo sufficient follow-up; thus, laboratories are encouraged to communicate the possibility of HSIL or other significant abnormality in their reports.

Reporting and management of histologic glandular lesions

Adenocarcinoma in situ (AIS)
The Society of Gynecologic Oncology (SGO) recently completed guidelines on the management of AIS and the 2019 ASCCP risk-based management consensus guidelines for glandular lesions are harmonized with the SGO guidelines.\textsuperscript{46} The only additional SGO-specific recommendation for management of an abnormal HPV-based screen result is that endocervical sampling is acceptable for any patient who tests positive for HPV 18 because of the high rate of HPV 18-positive AIS.

A diagnostic excisional procedure is recommended for all patients with a diagnosis of AIS on cervical biopsy to rule out invasive adenocarcinoma, even if hysterectomy is planned. The guidelines do not specifically recommend cold knife conization over loop electrocautery excisional procedure (LEEP), but do state that excisional procedures should aim to remove an intact specimen for evaluation of margin status by the pathologists. Hence, performance of a LEEP followed by a “top hat” endocervical excision is not acceptable. Ultimately, hysterectomy remains the preferred management for all patients with histologic AIS, but a fertility-sparing procedure is considered acceptable in select patients. Hysterectomy is preferred for AIS because it is often located within the endocervical canal, and is difficult to recognize on colposcopy, thus making decision on the extent of excision difficult. AIS is also known to have a higher risk of being multifocal, so negative margins on an
excisional specimen do not ensure complete excision of disease. After hysterectomy, surveillance is recommended per the 2019 ASCCP guidelines for treated CIN 2+.15,46

Newer glandular entities
Clinical providers are often not familiar with newer histologic entities, such as the stratified mucin-producing intraepithelial lesion (SMILE), a form of intraepithelial lesion distinct from conventional squamous and glandular counterparts.47 An invasive form of SMILE has recently been reported.48,49 In the largest retrospective cytology study to date, Backhouse et al found that almost 90% of SMILE were interpreted as squamous lesions on initial cytology and only about 10% were classified as glandular lesions.50 Pathologists should consider providing a short description of the significance of newer entities in their reports and/or discussing with the clinical provider the significance of newer terminology, so as to provide guidance on management.

Pregnancy-related considerations
Pregnancy is a special management consideration that weighs the risk of harm to fetus and mother against the risk of missing cancer. Although these should not be ignored, the evidence suggests that diagnostic procedures are safe in the hands of experienced colposcopists and that the rates of CIN 3+ progression are not higher in the pregnant woman.51 The goal of colposcopy during pregnancy is to exclude cancer, rather than to find high-grade disease. Adding to the complexity, however, are the challenges that colposcopists have detecting and recognizing lesions on the pregnant cervix. Visual recognition of CIN 3+ may be compromised by physiologic changes and cervical hyperemia of pregnancy, leading to underdiagnosis of CIN 3+. Colposcopically, decidual change on the cervix can be overinterpreted as suspicious for cancer, and cellular changes such as Arias-Stella changes and decidua maybe interpreted as AGC, ASC-H, ASC-US, and LSIL by cytologists.26

Social issues also complicate treatment timing decisions. Patients may not attend postpartum follow-up because of employment issues and demands of a new baby; in addition, an estimated 11% of women lose their health insurance in the postpartum period.52 For these reasons, some clinicians may determine that prepartum diagnostic procedures are appropriate. The 2019 ASCCP guidelines recommend the same clinical action thresholds for management, surveillance, and colposcopy for pregnant women as for non-pregnant women, except that endocervical curettage, endometrial biopsies, and expedited treatment (eg, conization or LEEP without histologic confirmation) are not recommended. Although colposcopy, biopsy, and cytology with age-appropriate HPV testing should occur during pregnancy, treatment for CIN 2/3 is postponed until postpartum. Cervical biopsies may be obtained during pregnancy in cases of suspected invasive carcinoma.

Quality assurance: cytology-histology correlation and more
Converting to a risk-based management strategy with primary HPV screening, especially in vaccinated individuals, introduces potential future diagnostic problems. HPV tests are highly sensitive but not specific—they identify the presence of the virus but don’t define the disease. Colposcopy has its own set of limitations and risks, as will the use of cervical cytology and biopsy as diagnostic tests. That is why the 2019 ASCCP guidelines recommend that all positive HPV tests receive cytology as a reflex test to aid with management, and recommend collection of a cytology sample at the time of colposcopy if reflex cytology from the screening sample is not possible.

Pathologist concordance with a particular cervical cytology or cervical biopsy result varies depending on the lesion. Overall, diagnostic reproducibility between general surgical pathologists and gynecologic pathologists regarding cervical biopsies are typically moderate, at best.53-55 Pathologists are most concordant at the far ends of the diagnostic spectrum (negative and CIN 3)56 but agree less often regarding CIN 1 and CIN 2.57 The histologic distinction between CIN 1 and CIN 2 is often nebulous, presenting a problem if the trigger for treatment is at CIN 2. The ASCUS-LSIL Triage Study demonstrated that diagnostic concordance for CIN 1/LSIL was greater for cervical cytology than for cervical biopsies.58 The diagnosis of CIN 1 on biopsies has only fair to moderate concordance. In a study by Basu et al, the lowest agreement among pathologists was for CIN 1 whereas the highest was for squamous cell carcinoma.56 Even when pathologists use a 3-tiered system of cervical biopsy interpretation, such as negative, LSIL, and HSIL, there is still difficulty separating normal cervical biopsies from LSIL and separating LSIL from HSIL.58 There is a tendency for pathologists to overcall normal epithelium as CIN 1 and to overcall CIN 1 as CIN 2.3,54 The use of p16 may be helpful in equivocal cases, as discussed above. A negative p16 is useful in excluding HSIL, but p16 positivity only indicates an HPV-associated lesion and does not differentiate HSIL from LSIL. Although cytology is more accurate than histology in diagnosing CIN 1, the distinction of CIN 2 from CIN 3 is problematic in both tissue and cytology, with poor to moderate reproducibility in both.57,59 In educational settings, such as the College of American Pathologists Educational Glass Slide Program, both pathologists and cytotechnologists are more likely to undercall HSIL as LSIL on cytology than the reverse.56 Papanicolaou tests with mixed LSIL and HSIL do not perform well in these circumstances either.61 These diagnostic tendencies may continue to confound management algorithms, skew future
data, and serve as a reminder that clinical management should not be solely reliant on test results alone.

The 2019 ASCCP guidelines continue to encourage clinical provider and pathologist interaction with review of the specimens when there is a significant cytologic, histologic, or colposcopic discrepancy. For example: “when CIN 2+ is not identified histologically after an ASC-H or HSIL cytology result, it is acceptable to review the cytologic, histologic, and colposcopic findings; if the review yields a revised interpretation, management should follow guidelines for the revised diagnosis.” Optimal practice includes real-time cytologic-histologic correlation with efforts to resolve discrepancies, such as obtaining deeper levels and/or performing p16 staining on histology if the preceding cytology result is of a higher-grade lesion. Making note of the review of cases, action taken, and resolution in the histology report is ideal.

Diagnostic competence in surgical pathology and cytopathology depends upon exposure; that is, seeing an appropriate volume of cases and making comparisons with other clinical and laboratory data. As a quality assurance metric, cytologic-histologic correlation plays a vital role in providing feedback on accuracy and “fine tuning” of diagnostic criteria. The opportunity for cytologic-histologic correlation will decrease as cytology is used primarily as triage or follow-up test. In an era where fewer cytology tests are reviewed, and most of these are HPV-positive, the proportion of atypical cytology results is likely to increase because of reviewer bias that knowledge of HPV status introduces. The consequence may be more colposcopy referrals for women without disease.

The complex and highly regulated quality environment federally mandated by the Clinical Laboratory Improvement Amendments (CLIA) for the practice of gynecologic cytopathology resulted in a robust system of quality assurance that is fully-developed and highly successful. Most surgical pathology quality systems in place are not as robust, and this may affect the interpretation of cervical biopsies. For example, a minimum of 2 individuals (a cytotechnologist and a pathologist) usually reviews all abnormal cervical cytology cases, but a single pathologist may solely diagnose cervical biopsies. There is no required peer review, interpreter monitoring, or correlation with outcomes for cervical biopsies similar to that mandated for cervical cytology. With higher stakes for missing a cervical abnormality due to extended screening intervals and follow-up, it may become a future best practice that more cervical biopsies receive a second review.

Conclusion

The landscape of cervical cancer prevention is rapidly changing. HPV immunization is reducing infections caused by targeted high-risk genotypes in vaccinated populations and reducing the incidence of high-grade lesions, caused by these types, whose morphology is most familiar to pathologists. Molecular approaches to screening with HPV tests are supplanting our familiar, but decades-old, morphology-based screening approach, the Papanicolaou test. New biomarkers, such as dual staining for screening, have recently received FDA approval. Additional molecular tests, such as extended HPV genotyping and type-persistence data from HPV tests that include partial and extended genotyping, need to be incorporated into the assessment of progression risk for those with histologic HSIL. New molecular tests, such as viral and host methylation, are on the horizon and promise to provide more objective and precise assessments of the risk for true precancer. In the future, pathologists may also have additional biomarkers for histopathology that more accurately reflect a lesion’s true risk for cancer progression. The clinical management guidelines will continue to evolve and, hopefully, more accurately balance the benefits and potential harms of cervical cancer prevention efforts. The World Health Organization has set targets for the elimination and eradication of cervical cancer. As laboratory workers, we must keep abreast of these new developments and guidelines and be prepared to proactively participate in and facilitate multidisciplinary secondary prevention of cervical cancer.

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