Human Epidermal Growth Factor Receptor 2 Testing in Breast Cancer

American Society of Clinical Oncology/College of American Pathologists Clinical Practice Guideline Focused Update

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• Purpose.—To update key recommendations of the American Society of Clinical Oncology (ASCO)/College of American Pathologists (CAP) human epidermal growth factor receptor 2 (HER2) testing in breast cancer guideline.

Methods.—Based on the signals approach, an Expert Panel reviewed published literature and research survey results on the observed frequency of less common in situ hybridization (ISH) patterns to update the recommendations.

Recommendations.—Two recommendations addressed via correspondence in 2015 are included. First, immunohistochemistry (IHC) 2+ is defined as invasive breast cancer with weak to moderate complete membrane staining observed in >10% of tumor cells. Second, if the initial HER2 test result in a core needle biopsy specimen of a primary breast cancer is negative, a new HER2 test may (not “must”) be ordered on the excision specimen based on specific clinical criteria. The HER2 testing algorithm for breast cancer is updated to address the recommended workup for less common clinical scenarios (approximately 5% of cases) observed when using a dual-probe ISH assay. These scenarios are described as ISH group 2 (HER2/chromosome enumeration probe 17 [CEP17] ratio >2.0; average HER2 copy number <4.0 signals per cell), ISH group 3 (HER2/CEP17 ratio <2.0; average HER2 copy number >6.0 signals per cell), and ISH group 4 (HER2/CEP17 ratio <2.0; average HER2 copy number >4.0 and <6.0 signals per cell). The diagnostic approach includes more rigorous interpretation criteria for IHC and requires concomitant IHC review for dual-probe ISH groups 2 to 4 to arrive at the most accurate HER2 status designation (positive or negative) based on combined interpretation of the ISH and IHC assays. The Expert Panel recommends that laboratories using single-probe IHC assays include concomitant IHC review as part of the interpretation of all single-probe IHC assay results.

THE BOTTOM LINE


Guideline Questions
What is the most appropriate definition for immunohistochemistry (IHC) 2+ (IHC equivocal)? Must human epidermal growth factor receptor 2 (HER2) testing be repeated on a surgical specimen if there was an initially negative test result on core biopsy? What is the optimal algorithm for less common patterns observed when performing dual-probe in situ hybridization (ISH) testing in breast cancer?

Target Population
Patients with breast cancer.

Target Audience
Medical oncologists, pathologists, surgeons, and radiation oncologists.

Methods
An Expert Panel was convened to develop updated clinical practice guideline recommendations based on a systematic review of the medical literature.

Focused Update Recommendations

1. In the revised Figure 1, the revised definition of IHC 2+ (equivocal) is invasive breast cancer with “weak to moderate complete membrane staining observed in >10% of tumor cells.”

2. In the revised Table 2, it is now stated that, on the basis of some criteria (including a tumor grade 3), “If the initial HER2 test result in a core needle biopsy specimen of a primary breast cancer is negative, a new HER2 test may be ordered on the excision specimen…”

3. If a case has an HER2/chromosome enumeration probe 17 (CEP17) ratio of ≥2.0 but the average HER2 signals per cell is <4.0, a definitive diagnosis will be rendered based on additional workup. If not already assessed by the institution or laboratory performing the ISH test, IHC testing for HER2 should be performed using sections from the same tissue sample used for ISH, and the slides from both ISH and IHC should be reviewed together to guide the selection of areas to score by ISH (local practice considerations will dictate the best procedure to accomplish this concomitant review):
   a. If the IHC result is 3+, diagnosis is HER2 positive.
   b. If the IHC result is 2+, recount ISH by having an additional observer, blinded to previous ISH results, count at least 20 cells that include the area of invasive cancer with IHC 2+ staining:
      • If reviewing the count by the additional observer changes the result into another ISH category, the result should be adjudicated per internal procedures to define the final category.
      • If the count remains an average of <4.0 HER2 signals per cell and the HER2/CEP17 ratio is ≥2.0, diagnosis is HER2 negative with a comment. (Please note: Refer to text for specific comments about recommendations listed as 3b, 3c, 4c, and 5b).
   c. If the IHC result is 0 or 1+, diagnosis is HER2 negative with a comment. (Please note: Refer to text for specific comments about recommendations listed as 3b, 3c, 4c, and 5b).

4. If a case has an average of ≥6.0 HER2 signals per cell with an HER2/CEP17 ratio of <2.0, formerly diagnosed as ISH positive for HER2, a definitive diagnosis will be rendered based on additional workup. If not already assessed by the institution or laboratory performing the ISH test, IHC testing for HER2 should be performed using sections from the same tissue sample used for ISH, and the slides from both ISH and IHC should be reviewed together to guide the selection of areas to score by ISH (local practice considerations will dictate the best procedure to accomplish this concomitant review):
   a. If the IHC result is 3+, diagnosis is HER2 positive.
   b. If the IHC result is 2+, recount ISH by having an additional observer, blinded to previous ISH results, count at least 20 cells that include the area of invasion with IHC 2+ staining:
      • If reviewing the count by the additional observer changes the result into another ISH category, the result should be adjudicated per internal procedures to define the final category.
      • If the HER2/CEP17 ratio remains <2.0 with ≥6.0 HER2 signals per cell, diagnosis is HER2 positive.
   c. If the IHC result is 0 or 1+, diagnosis is HER2 negative with a comment. (Please note: Refer to text for specific comments about recommendations listed as 3b, 3c, 4c, 5b, and 5c).

5. If the case has an average HER2 signals per tumor cell of ≥4.0 and <6.0 and the HER2/CEP17 ratio is <2.0, formerly diagnosed as ISH equivocal for HER2, a definitive diagnosis will be rendered based on additional workup. If not already assessed by the institution or laboratory performing the ISH test, IHC testing for HER2 should be performed using sections from the same tissue sample used for ISH, and the slides from both ISH and IHC should be reviewed together to guide the selection of areas to score by ISH (local practice considerations will dictate the best procedure to accomplish this concomitant review):
   a. If the IHC result is 3+, diagnosis is HER2 positive.
   b. If the IHC result is 2+, recount ISH by having an additional observer, blinded to previous ISH results, count at least 20 cells that include the area of invasion with IHC 2+ staining:
      • If reviewing the count by the additional observer changes the result into another ISH category, the result should be adjudicated per internal procedures to define the final category.
      • If the count remains an average of ≥4.0 and <6.0 HER2 signals per cell with an HER2/CEP17 ratio of <2.0, diagnosis is HER2 negative with a comment. (Please note: Refer to text for specific comments about recommendations listed as 3b, 3c, 4c, 5b, and 5c).
   c. If the IHC result is 0 or 1+, diagnosis is HER2 negative with a comment. (Please note: Refer to text for specific comments about recommendations listed as 3b, 3c, 4c, 5b, and 5c).

Note: In Figure 2, a new footnote states that the Expert Panel recommends that concomitant IHC review become part of the interpretation of single-probe ISH results and that the Expert Panel preferentially recommends the use of dual-probe instead of single-probe ISH assays.

Refer to Table 1 for the full list of recommendations.
Table 1. Summary of All Recommendations (Original Recommendations and Focused Update Recommendations)

<table>
<thead>
<tr>
<th>Topic</th>
<th>2013 Recommendations</th>
<th>2018 Focused Update Recommendations</th>
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</thead>
<tbody>
<tr>
<td>Specimens to be tested</td>
<td>All newly diagnosed patients with breast cancer must have an HER2 test performed. Patients who then develop metastatic disease must have an HER2 test performed in a metastatic site, if tissue sample is available.</td>
<td>No change.</td>
</tr>
</tbody>
</table>
| Optimal algorithm for HER2 testing | Must report HER2 test result as positive for HER2 if: IHC 3+ based on circumferential membrane staining that is complete, intense ISH positive based on: Single-probe average HER2 copy number ≥ 6.0 signals/cell Dual-probe HER2/CEP17 ratio of ≥ 2.0, with an average HER2 copy number ≥ 4.0 signals/cell Dual-probe HER2/CEP17 ratio of ≥ 2.0, with an average HER2 copy number < 4.0; Dual-probe HER2/CEP17 ratio of < 2.0, with an average HER2 copy number ≥ 6.0 signals/cell | 1. In the revised Figure 1, the revised definition of IHC 2+ (equivocal) is invasive breast cancer with “weak to moderate complete membrane staining observed in > 10% of tumor cells.”
2. In the revised Table 2, it is now stated that, on the basis of some criteria (including a tumor grade 3), “If the initial HER2 test result in a core needle biopsy specimen of a primary breast cancer is negative, a new HER2 test may be ordered on the excision specimen.”
3. If a case has an HER2/CEP17 ratio of ≥ 2.0 but the average HER2 signals/cell is < 4.0, a definitive diagnosis will be rendered based on additional workup. If not already assessed by the institution or laboratory performing the ISH test, IHC testing for HER2 should be performed using sections from the same tissue sample used for ISH, and the slides from both ISH and IHC should be reviewed together to define the selection of areas to score by ISH (local practice considerations will dictate the best procedure to accomplish this concomitant assessment):
   a. If the IHC result is 3+, diagnosis is HER2 positive
   b. If the IHC result is 2+, recount ISH by having an additional observer blinded to previous ISH results, count at least 20 cells that include the area of invasive cancer with IHC 2+ staining:
      1) If reviewing the count by the additional observer changes the result into another ISH category, the result should be adjudicated per internal procedures to define the final category
      2) If the count remains an average of < 4.0 HER2 signals/cell and the HER2/CEP17 ratio is ≥ 2.0, diagnosis is HER2 negative with a comment*
   c. If the IHC result is 0 or 1+, diagnosis is HER2 negative with a comment*
4. If a case has an average of ≥ 6.0 HER2 signals/cell with an HER2/CEP17 ratio of < 2.0, formerly diagnosed as ISH positive for HER2, a definitive diagnosis will be rendered based on additional workup. If not already assessed by the institution or laboratory performing the ISH test, IHC testing for HER2 should be performed using sections from the same tissue sample used for ISH, and the slides from both ISH and IHC should be reviewed together to guide the selection of areas to score by ISH (local practice considerations will dictate the best procedure to accomplish this concomitant review):
   a. If the IHC result is 3+, diagnosis is HER2 positive
   b. If the IHC result is 2+, recount ISH by having an additional observer blinded to previous ISH results, count at least 20 cells that include the area of invasion with IHC 2+ staining:
      1) If reviewing the count by the additional observer changes the result into another ISH category, the result should be adjudicated per internal procedures to define the final category
      2) If the HER2/CEP17 ratio remains < 2.0 with ≥ 6.0 HER2 signals/cell, diagnosis is HER2 positive
   c. If the IHC result is 0 or 1+, diagnosis is HER2 negative with a comment*
5. If the case has an average HER2 signal/tumor cell of ≥ 4.0 and < 6.0 and the HER2/CEP17 ratio is < 2.0, formerly diagnosed as ISH equivocal for HER2, a definitive diagnosis will be rendered based on additional workup. If not already assessed by the institution or laboratory performing the ISH test, IHC testing for HER2 should be performed using sections from the same tissue sample used for ISH, and the slides from both ISH and IHC should be reviewed together to guide the selection of areas to score by ISH (local practice considerations will dictate the best procedure to accomplish this concomitant review):
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      1) If reviewing the count by the additional observer changes the result into another ISH category, the result should be adjudicated per internal procedures to define the final category
      2) If the HER2/CEP17 ratio remains < 2.0 with ≥ 6.0 HER2 signals/cell, diagnosis is HER2 positive
   c. If the IHC result is 0 or 1+, diagnosis is HER2 negative with a comment*
<table>
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<th>Topic</th>
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</tr>
</thead>
<tbody>
<tr>
<td>ISH rejection criteria</td>
<td>Test is rejected and repeated if: Controls are not as expected. Observer cannot find</td>
<td>The pathologist should scan the entire ISH slide before counting at least 20 cells or use IHC to</td>
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<td></td>
<td>and count at least two areas of invasive tumor. &gt;25% of signals are unscorable due</td>
<td>define the areas of potential HER2 amplification. If there is a second population of contiguous cells</td>
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<td></td>
<td>to weak signals &gt;10% of signals occur over cytoplasm Nuclear resolution is poor</td>
<td>with increased HER2 signals/cell and this cell population consists of &gt;10% of tumor cells on the</td>
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<td>Autofluorescence is strong</td>
<td>slide (defined by image analysis or visual estimation of the ISH or IHC slide), a separate counting</td>
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<td></td>
<td>Report HER2 test result as Indeterminate as per parameters described</td>
<td>of at least 20 nonoverlapping cells must also be performed within this cell population and reported.</td>
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<tr>
<td>ISH interpretation</td>
<td>The pathologist should scan the entire ISH slide before counting at least 20 cells or</td>
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<td></td>
<td>or use IHC to define the areas of potential HER2 amplification.</td>
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<td></td>
<td>If there is a second population of cells with increased HER2 signals/cell and this</td>
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<td></td>
<td>cell population consists of &gt;10% of tumor cells on the slide (defined by image analysis</td>
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<td></td>
<td>or visual estimation of the ISH or IHC slide), a separate counting of at least 20</td>
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<td></td>
<td>nonoverlapping cells must also be performed within this cell population and reported.</td>
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<td></td>
<td>For brightfield ISH, counting requires comparison between patterns in normal breast</td>
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<td>and tumor cells because artifactual patterns may be seen that are difficult to interpret. If tumor cell</td>
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<td>pattern is neither normal nor clearly amplified, test should be submitted for expert</td>
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<td>opinion.</td>
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<td>Acceptable (IHC and ISH)</td>
<td>Should preferentially use an FDA-approved IHC, brightfield ISH, or FISH assay.</td>
<td>No change</td>
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<td>tests</td>
<td>No change</td>
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<tr>
<td>IHC rejection criteria</td>
<td>Test is rejected, and repeated or tested by FISH if: Controls are not as expected</td>
<td>No change</td>
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<td></td>
<td>Artifacts involve most of sample</td>
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<td></td>
<td>Sample has strong membrane staining of normal breast ducts (internal controls)</td>
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<tr>
<td>IHC interpretation criteria</td>
<td>Should interpret IHC test using a threshold of &gt;10% of tumor cells that must show</td>
<td>No change</td>
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<td>homogeneous, dark circumferential (chicken wire) pattern to call result 3+, HER2</td>
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<td>positive.</td>
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<td>Reporting requirements</td>
<td>Report must include guideline-detailed elements except for changes to reporting</td>
<td>No change</td>
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<td>for all assay types</td>
<td>requirement and algorithms defined in this table.</td>
<td></td>
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<tr>
<td>Optimal tissue handling</td>
<td>Time from tissue acquisition to fixation should be as short as possible; samples for HER2</td>
<td>No change</td>
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<tr>
<td>requirements</td>
<td>testing are fixed in 10% neutral buffered formalin for 6-72 hours; cytology specimens</td>
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<td>must be fixed in formalin. Samples should be sliced at 5- to 10-mm intervals after</td>
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<td>appropriate gross inspection and margin designation and placed in a sufficient volume</td>
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<td>of neutral buffered formalin. Any exceptions to this process must be included in the</td>
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<td>report.</td>
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<td>Topic</td>
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<tr>
<td>Optimal tissue sectioning requirements</td>
<td>Sections should ideally not be used for HER2 testing if cut &gt;6 weeks earlier; this may vary with primary fixation or storage conditions</td>
<td>No change</td>
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<td>Optimal internal validation procedure</td>
<td>Validation of test must be performed before test is offered</td>
<td>No change</td>
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<tr>
<td>Optimal initial test validation</td>
<td>Laboratories performing these tests should be following all accreditation requirements, one of which is initial testing validation. The laboratory should ensure that initial validation conforms to the published 2010 ASCO/CAP recommendations for IHC testing of ER and PgR guideline validation requirements with 20 negative and 20 positive for FDA-approved assays and 40 negative and 40 positive for LDTs. This requirement does not apply to assays that were previously validated in conformance with the 2007 ASCO/CAP HER2 testing guideline, and those who routinely participate in external proficiency testing for HER2 tests, such as the program offered by CAP. Laboratories are responsible for ensuring the reliability and accuracy of their testing results, by compliance with accreditation and proficiency testing requirements for HER2 testing assays. Specific concordance requirements are not required.</td>
<td>No change</td>
</tr>
<tr>
<td>Optimal monitoring of test concordance between methods</td>
<td>See text below, under optimal laboratory accreditation</td>
<td>No change</td>
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<tr>
<td>Optimal internal QA procedures</td>
<td>Should review and document external and internal controls with each test and each batch of tests Ongoing quality control and equipment maintenance Initial and ongoing laboratory personnel training and competency assessment Use of standardized operating procedures, including routine use of control materials Revalidation of procedure if changed Ongoing competency assessment and documentation of the actions taken as a part of the laboratory record</td>
<td>No change</td>
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<tr>
<td>Optimal external proficiency assessment</td>
<td>Participation in and successful completion of external proficiency testing program with at least two testing events (mailings) a year Satisfactory performance requires at least 90% correct responses on graded challenges for either test Unsatisfactory performance will require laboratory to respond according to accreditation agency program requirements</td>
<td>No change</td>
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</table>
testing and the clinical utility of HER2 as a predictive biomarker for potential responsiveness to therapies targeting the HER2 protein.

Activating mutations of the tyrosine kinase and extracellular domains of HER2 in the absence of amplification or overexpression offer an alternative and much less common mechanism for HER2-targeted therapy that is being explored in clinical trials of small molecule kinase inhibitors. Data from NRG trial B-47 (ClinicalTrials.gov identifier: NCT01275677) confirmed the lack of benefit from adjuvant trastuzumab for patients whose tumors lack gene amplification and are immunohistochemistry (IHC) 1+ or 2+. Consequently, HER2 gene amplification assessed by in situ hybridization (ISH) or protein overexpression assessed by IHC remains the primary predictor of responsiveness to HER2-targeted therapies in breast cancer.

Greater communication among health care providers (especially pathologists and oncologists) and appropriate infrastructure support for specimen handling and laboratory facilities led to observed improvements in the analytic performance and accuracy of HER2 testing. Greater clinical experience with the efficacy and safety of HER2-targeted therapies, and a meaningful reduction in the high frequency of false-positive HER2 test results previously observed, led the 2013 Expert Panel to provide additional guidance regarding less common clinical scenarios to allow greater discrimination between positive and negative results.

Since 2013, several laboratory and clinical investigators have reported on the practical implications of the 2013 Guideline Update and the observed frequency of equivocal cases.

The HER2 testing Guideline Expert Panel has identified five clinical questions that form the core of this 2018 Focused Update. Two of them (Clinical Questions 1 and 2) were addressed in a previous correspondence by the panel that was published in the *Journal of Clinical Oncology* (JCO) in 2015, and they are included here in the forms of Figure 1 (algorithm for IHC testing) and Table 2 (histopathologic features suggestive of possible test discordance), both revised from the 2013 guideline. Figure 2 is an algorithm for single-probe ISH testing and includes a new footnote with a recommendation that concomitant IHC review become part of the interpretation of single-probe ISH results. Clinical Questions 3, 4, and 5 address less common patterns observed when performing dual-probe ISH testing and are graphically summarized in Figures 3 to 6 (algorithm for dual-probe ISH testing), also revised.

A new Table 3 describes the patterns of HER2 ISH testing using a dual-probe assay and lists the clear effect of the underlying distribution of HER2 IHC test results on the frequency of less common patterns of ISH (hereafter called groups 2, 3, and 4). In the population at large, approximately 95% of tumors tested for HER2 by dual-
probe ISH will consist of group 1 (HER2 positive) and group 5 (HER2 negative). It is expected that approximately 5% of cases tested by ISH will fall into groups 2, 3, or 4, and available clinical outcome data from related clinical trials, albeit of limited statistical power, have allowed the Expert Panel to more carefully define the expected prognostic and predictive behavior of these cases.

Most importantly, after careful consideration of the available evidence and expert opinions, the Expert Panel revised the diagnostic approach to groups 2 to 4 to include more rigorous interpretation criteria for dual-probe ISH testing and to require concomitant IHC review, to arrive at the most accurate HER2 status designation (positive or negative) based on the combined interpretation of the ISH and IHC assays. The Expert Panel recommends that such concomitant review be performed in the same institution to ensure parallel interpretation and quality of the two assays.

Although the main focus was to clarify the less common test results observed with the two-probe ISH assays, the recommendations affect the users of single-probe ISH assays. Therefore, the Expert Panel now recommends that concomitant IHC review become part of the interpretation of single-probe ISH results to allow the most accurate HER2 designation (Figure 2). The Expert Panel also preferentially recommends the use of dual-probe instead of single-probe ISH assays, although it recognizes that several single-probe ISH assays have regulatory approval in many parts of the world.

GUIDELINE QUESTIONS

This 2018 Focused Update addresses five clinical questions raised after the publication of the 2013 Guideline Update:

**Clinical Question 1**

What is the most appropriate definition for IHC 2+ (IHC equivocal)?

**Clinical Question 2**

Must HER2 testing be repeated on a surgical specimen if initially negative test on core biopsy?

**Clinical Question 3**

Should invasive cancers with an HER2/chromosome enumeration probe 17 (CEP17) ratio of ≥2.0 but an average HER2 copy number of <4.0 signals per cell be considered ISH positive?

**Clinical Question 4**

Should invasive cancers with an average HER2 copy number of ≥6.0 signals per cell but a HER2/CEP17 ratio of <2.0 be considered ISH positive?

**Clinical Question 5**

What is the appropriate diagnostic workup for invasive cancers with an average HER2 copy number of ≥4.0 but <6.0 signals per cell and an HER2/CEP17 ratio of <2.0, and initially deemed to have an equivocal HER2 ISH test result?
Table 2. Histopathologic Features Suggestive of Possible Human Epidermal Growth Factor Receptor 2 (HER2) Test Discordance*

<table>
<thead>
<tr>
<th>Criteria to Consider†</th>
<th>A new HER2 test should not be ordered if the following histopathologic findings occur and the initial HER2 test was negative:</th>
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<tbody>
<tr>
<td></td>
<td>Histologic grade 1 carcinoma of the following types: Infiltrating ductal or lobular carcinoma, ER and PgR positive Tubular (at least 90% pure) Mucinous (at least 90% pure) Cribriform (at least 90% pure) Adenoid cystic carcinoma (90% pure) and often triple negative</td>
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<tr>
<td></td>
<td>Similarly, a new HER2 test should be ordered if the following histopathologic findings occur and the initial HER2 test was positive:</td>
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<tr>
<td></td>
<td>Histologic grade 1 carcinoma of the following types: Infiltrating ductal or lobular carcinoma, ER and PgR positive Tubular (at least 90% pure) Mucinous (at least 90% pure) Cribriform (at least 90% pure) Adenoid cystic carcinoma (90% pure) and often triple negative</td>
</tr>
<tr>
<td></td>
<td>If the initial HER2 test result in a core needle biopsy specimen of a primary breast cancer is negative, a new HER2 test may be ordered on the excision specimen if one of the following is observed:</td>
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<td>Tumor is grade 3</td>
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<td>Amount of invasive tumor in the core biopsy specimen is small</td>
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<td>Resection specimen contains high-grade carcinoma that is morphologically distinct from that in the core</td>
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<td>Core biopsy result is equivocal for HER2 after testing by both ISH and IHC</td>
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<td></td>
<td>There is doubt about the handling of the core biopsy specimen (long ischemic time, short time in fixative, different fixative), or the test is suspected by the pathologist to be negative on the basis of testing error</td>
</tr>
</tbody>
</table>

Abbreviations: ER, estrogen receptor; IHC, immunohistochemistry; ISH, in situ hybridization; PgR, progesterone receptor.
* Adapted from the 2013 ASCO/CAP HER2 Testing Guideline.† Criteria to consider if there are concerns regarding discordance with the 2013 ASCO/CAP HER2 Testing Guideline published in JCO13 in 2015 in response to correspondence by Rakha et al,17 and this Focused Update contains a revised Table 1 (Clinical Question 1) and a revised Table 2 (Clinical Question 2).

Clinical Questions 1 and 2 were formally addressed in correspondence from the ASCO/CAP HER2 testing Expert Panel published in JCO13 in 2015 in response to correspondence by Rakha et al,17 and this Focused Update contains a revised Figure 1 (Clinical Question 1) and a revised Table 2 (Clinical Question 2).

Clinical Questions 3, 4, and 5 regarding dual-probe (dual-signal) ISH testing (Figures 3 to 6) were addressed by the Expert Panel in a meeting at ASCO headquarters on November 28 and 29, 2016, and in subsequent conference calls and electronic communications. Figure 2 (single-probe ISH) from the 2013 Guideline Update includes a new footnote with a recommendation that concomitant IHC review become part of the interpretation of single-probe ISH results. Table 1 contains a summary of the recommendations of the 2013 Guideline Update and the 2018 Focused Update. Figure 7, describing the number of laboratories participating in predictive marker proficiency testing for HER2, has been updated.

METHODS

Guideline Update Process

This systematic review-based guideline product was developed by a multidisciplinary Expert Panel, which included a patient representative and ASCO guidelines staff with health research methodology expertise. PubMed and the Cochrane Library were searched for randomized controlled trials, systematic reviews, meta-analyses, and clinical practice guidelines for the period from January 1, 2013, through May 11, 2017. The disease and intervention search terms were those that were used for the 2013 Guideline Update. The updated search was guided by the signals18 approach that is designed to identify only new, potentially practice-changing data (signals) that might translate into revised practice recommendations. The approach relies on targeted routine literature searching and the expertise of ASCO Expert Panel members to help identify potential signals. Additional information about the literature search strategy string and results, as well as a discussion of the ASCO signals approach to guideline updating, are available in the 2018 Data Supplement and 2018 Methodology Supplement, respectively (see supplemental digital content at www.archivesofpathology.org in the November 2018 table of contents). A QUOROM diagram of the updated search and the clinical questions are provided (Data Supplement). In addition to the literature search, a research survey was distributed to gather additional real-world data from laboratories from before and after implementation of the ASCO/CAP HER2 Testing in Breast Cancer 2013 Update. Additional information regarding this survey process is available in the Data Supplement.

The Expert Panel met during a 2-day in-person meeting in November 2016 to consider the evidence for each of the recommendations contained in this 2018 Focused Update. Laboratories that shared with the Expert Panel their clinical experience with HER2 testing since the publication of the 2013 Guideline Update participated in the open session of the meeting. The guideline was circulated in draft form to the Expert Panel. Draft recommendations were released to the public for an open comment period between May 22 and June 19, 2017. ASCO’s Clinical Practice Guidelines Committee reviewed and approved the final document. For CAP, an independent review panel was assembled to review and approve the guideline. The independent review panel was masked to the Expert Panel and was vetted through the conflict of interest process.

Only recommendations relating to the updated clinical questions have changed. The Data Supplement provides clinical questions corresponding to all recommendations from the 2013 Guideline Update.

This ASCO/CAP Clinical Practice Guideline Focused Update provides select recommendations and a comprehensive discussion of the relevant literature from January 1, 2013, to May 11, 2017, for these specific recommendations. The full guideline, which this revision applies to, and additional information are available at www.asco.org/breast-cancer-guidelines. The complete list of recommendations, including the updated recommendations, is in Table 1. All funding for the administration of the project was provided by ASCO and CAP.

Guideline Disclaimer

The clinical practice guidelines and other guidance published herein are provided by the American Society of Clinical Oncology Inc (“ASCO”) to assist providers in clinical decision-making. The information therein should not be relied upon as being complete or accurate, nor should it be considered as inclusive of all proper treatments or methods of care or as a statement of the standard of care. With the rapid development of scientific knowledge, new evidence may emerge between the time information is developed and when it is published or read. The information is not continually updated and may not reflect the most recent evidence. The information addresses only the topics specifically identified therein and is not applicable to other interventions, diseases, or stages of diseases. This information does not mandate any particular course of
medical care. Further, the information is not intended to substitute for the independent professional judgment of the treating provider, as the information does not account for individual variation among patients. Recommendations reflect high, moderate, or low confidence that the recommendation reflects the net effect of a given course of action. The use of words like “must,” “must not,” “should,” and “should not” indicates that a course of action is recommended or not recommended for either most or many patients, but there is latitude for the treating physician to select other courses of action in individual cases. In all cases, the selected course of action should be considered by the treating provider in the context of treating the individual patient. Use of the information is voluntary. ASCO

**Figure 2.** Algorithm for evaluation of human epidermal growth factor receptor 2 (HER2) gene amplification by in situ hybridization (ISH) assay of the invasive component of a breast cancer specimen using a single-signal (HER2 gene) assay (single-probe ISH). Note: The final reported results assume that there is no apparent histopathologic discordance observed by the pathologist. *It is recommended that concomitant immunohistochemistry (IHC) review become part of the interpretation of single-probe ISH results. The Expert Panel also preferentially recommends the use of dual-probe instead of single-probe ISH assays. †Using sections from the same tissue samples used for single-probe ISH, perform IHC (if not already performed) and/or dual-probe ISH. If IHC results are 2+ equivocal, it is recommended to also perform dual-probe ISH. ‡If initial assessment of dual-probe ISH is suggestive of groups 2, 3, or 4, follow the algorithm described in Figure 3.

**Figure 3.** Algorithm for evaluation of human epidermal growth factor receptor 2 (HER2) gene amplification by in situ hybridization (ISH) assay of the invasive component of a breast cancer specimen using a dual-signal (HER2 gene) assay (dual-probe ISH). Note: The final reported results assume that there is no apparent histopathologic discordance observed by the pathologist. Regarding groups 2, 3, and 4, if not already assessed by the institution or laboratory performing the ISH test, immunohistochemistry (IHC) testing for HER2 should be performed using sections from the same tissue sample used for ISH, and the slides from both ISH and IHC should be reviewed together to guide the selection of areas to score by ISH (local practice considerations will dictate the best procedure to accomplish this concomitant assessment). CEP17, chromosome enumeration probe 17.
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### Guideline and Conflicts of Interest

The Expert Panel was assembled in accordance with ASCO’s Conflict of Interest Policy Implementation for Clinical Practice Guidelines (“Policy,” found at www.asco.org/rwc) as agreed upon with CAP. All members of the Expert Panel completed ASCO’s disclosure form, which requires disclosure of financial and other

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**Figure 4. Clinical Question 3, group 2.**

*Evidence is limited on the efficacy of human epidermal growth factor receptor 2 (HER2)–targeted therapy in the small subset of cases with an HER2/chromosome enumeration probe 17 (CEP17) ratio ≥ 2.0 and an average HER2 copy number of <4.0 per cell. In the first generation of adjuvant trastuzumab trials, patients in this subgroup who were randomly assigned to the trastuzumab arm did not seem to derive an improvement in disease-free or overall survival, but there were too few such cases to draw definitive conclusions. Immunohistochemistry (IHC) expression for HER2 should be used to complement in situ hybridization (ISH) and define HER2 status. If the IHC result is not 3+ positive, it is recommended that the specimen be considered HER2 negative because of the low HER2 copy number by ISH and the lack of protein overexpression.*

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**Figure 5. Clinical Question 4, group 3.**

*There are insufficient data on the efficacy of human epidermal growth factor receptor 2 (HER2)–targeted therapy in cases with a HER2 ratio of ≥2.0 in the absence of protein overexpression because such patients were not eligible for the first generation of adjuvant trastuzumab clinical trials. When concurrent immunohistochemistry (IHC) results are negative (0 or 1+), it is recommended that the specimen be considered HER2 negative. CEP17, chromosome enumeration probe 17.*
Figure 6. Clinical Question 5, group 4. *It is uncertain whether patients with an average of ≥4.0 and <6.0 human epidermal growth factor receptor 2 (HER2) signals per cell and a HER2/chromosome enumeration probe 17 (CEP17) ratio of <2.0 benefit from HER2-targeted therapy in the absence of protein overexpression (immunohistochemistry [IHC] 3+). If the specimen test result is close to the in situ hybridization (ISH) ratio threshold for positive, there is a higher likelihood that repeat testing will result in different results by chance alone. Therefore, when IHC results are not 3+ positive, it is recommended that the sample be considered HER2 negative without additional testing on the same specimen.

Table 3. Distribution by Dual Fluorescent In Situ Hybridization (FISH) and Immunohistochemistry (IHC) Testing Results in Reported Data Sets*

<table>
<thead>
<tr>
<th>Laboratory</th>
<th>HERA Central Laboratory</th>
<th>BCIRG Central Laboratory</th>
<th>USC Breast Cancer Analysis Laboratory</th>
<th>Mayo Clinic Cytogenetics Laboratory</th>
<th>UK NEQAS 2009-2016†</th>
<th>Stanford/UCSF/UWMC16</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial Test Results</td>
<td>FISH distribution</td>
<td>IHC distribution</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No.</td>
<td>6018</td>
<td>10 468</td>
<td>7526</td>
<td>2851</td>
<td>11 116</td>
<td>8068</td>
</tr>
<tr>
<td>Group 1 ratio ≥2.0; HER2 ≥4.0</td>
<td>55.0 (6.0, 48.7; ≥4.0-6.0, 6.3)</td>
<td>Group 2 ratio ≥2.0; HER2 &lt;4.0</td>
<td>0.8</td>
<td>0.7</td>
<td>0.4</td>
<td>1.3</td>
</tr>
<tr>
<td>Group 3 ratio &lt;2.0; HER2 ≥6.0</td>
<td>0.4</td>
<td>0.5</td>
<td>0.6</td>
<td>3.0</td>
<td>1.1</td>
<td>0.8</td>
</tr>
<tr>
<td>Group 4 ratio &lt;2.0; HER2 ≥4.0 and &lt;6.0 (after alternative probe: pos, equivocal, neg)</td>
<td>1.9</td>
<td>4.1</td>
<td>4.6</td>
<td>14.2 (7.5, 5.5, 1.3)</td>
<td>7.6</td>
<td>5.2</td>
</tr>
<tr>
<td>Group 5 ratio &lt;2.0; HER2 &lt;4.0</td>
<td>41.9</td>
<td>53.9</td>
<td>76.7</td>
<td>69.6</td>
<td>73.4</td>
<td>78.8</td>
</tr>
<tr>
<td>IHC distribution</td>
<td>No.</td>
<td>3089</td>
<td>4331</td>
<td>7526</td>
<td>1922</td>
<td>11 116</td>
</tr>
<tr>
<td>0</td>
<td>IHC 0-1+, 2.0</td>
<td>54.5</td>
<td>51.7</td>
<td>2.4</td>
<td>0.3</td>
<td>IHC 0-1+, 38.1</td>
</tr>
<tr>
<td>1+ (including 0 or 1+)</td>
<td>—</td>
<td>9.4</td>
<td>31.0</td>
<td>8.0</td>
<td>1.8</td>
<td>—</td>
</tr>
<tr>
<td>2+ (including 1+/2+ or 2+/3+)‡</td>
<td>61.8</td>
<td>13.7</td>
<td>9.0</td>
<td>87.1‡</td>
<td>96.5‡</td>
<td>2+, 46.6</td>
</tr>
<tr>
<td>3+</td>
<td>36.2</td>
<td>22.4</td>
<td>8.4</td>
<td>2.5</td>
<td>1.3</td>
<td>3+, 15.3</td>
</tr>
</tbody>
</table>

Abbreviations: BCIRG, Breast Cancer International Research Group; HER2, human epidermal growth factor receptor 2; HERA, Herceptin Adjuvant trial; neg, negative; pos, positive; UCSF, University of California, San Francisco; UK NEQAS, United Kingdom National External Quality Assessment Service; USC, University of Southern California; UWMC, University of Washington Medical Center.

* Data are presented as % unless otherwise indicated.
† Andrew Dodson, personal communication, October 2016.
‡ IHC 1+ or 2+ and 2+ or 3+ were grouped together with IHC 2+. In each column for a specific laboratory or study, the top set of percentages describes the distribution of groups 1 to 5 results when tested using a dual-probe FISH assay, while the bottom set of percentages describes the distribution of IHC tests results of the samples submitted to that laboratory or study for dual-probe ISH testing and as described in each publication.
interests, including relationships with commercial entities that are reasonably likely to experience direct regulatory or commercial impact as a result of promulgation of the guideline. Categories for disclosure include employment; leadership; stock or other ownership; honoraria, consulting or advisory role; speaker’s bureau; research funding; patents, royalties, other intellectual property; expert testimony; travel, accommodations, expenses; and other relationships. In accordance with the Policy, the majority of the members of the Expert Panel did not disclose any relationships constituting a conflict under the Policy.

RECOMMENDATIONS

All recommendations regarding each of the five clinical questions are predicated on the assumption that the cases have been properly fixed, processed, and tested in a laboratory that follows ASCO/CAP HER2 testing guideline recommendations, especially those related to IHC and ISH interpretation and reporting.

In the 2013 Guideline Update, the workup of cases in the less common dual-probe ISH categories (groups 2 to 4) addressed in Clinical Questions 3, 4, and 5 (Figure 3) included only ISH. In this 2018 Focused Update, we recommend that these cases be worked up by considering both the IHC and the dual-probe ISH results together. Many publications since the 2013 Guideline Update have referenced the value of adjudicating ISH results in these uncommon categories using IHC.12,16,19–21 These tests should be performed on the same tissue sample using sections from the same block. Ideally, adjacent tissue levels from the same block should be tested and then reviewed together. If IHC has already been performed, it should be used to guide the selection of the areas to be counted during ISH such that areas with the strongest protein expression can be included in ISH scoring. This is common practice among laboratories performing both testing procedures. If the ISH laboratory only performs ISH, it is recommended that an adjacent section in the same block be assessed at a companion IHC laboratory, and then the slides from both ISH and IHC be reviewed together to guide the selection of areas to score by ISH. Local practice considerations will dictate the best procedure to accomplish this concomitant review.

Clinical Question 1

What is the most appropriate definition for IHC 2+ (IHC equivocal)?

2013 Recommendation.—IHC 2+ (equivocal) was defined in Figure 1 of the 2013 HER2 Testing Update as invasive breast cancer showing “circumferential membrane staining that is intense and within 10% of tumor cells.”

Revised 2018 Recommendation.—In the revised Figure 1, the revised definition of IHC 2+ (equivocal) is invasive breast cancer with “weak to moderate complete membrane staining observed in > 10% of tumor cells” (type: evidence based; evidence quality: high; strength of recommendation: strong).

Literature Review and Analysis.—IHC 2+ (equivocal) had been defined in the 2013 Guideline Update (2013 figure 1: Algorithm for evaluation of HER2 protein expression by IHC assay of the invasive component of a breast cancer specimen) as invasive breast cancer showing “circumferential membrane staining that is incomplete and/or weak/moderate and within > 10% of tumor cells or complete and circumferential membrane staining that is intense and within ≤ 10% of tumor cells.” However, many pathologists expressed concern that the terms “circumferential” and “incomplete” were confusing, could not be reconciled when used together in the IHC interpretation of HER2 expression, and could lead to many IHC 1+ (HER2-negative) tumors being called IHC 2+ (HER2 equivocal) and submitted for reflex testing.

In the same figure 1 of the 2013 Guideline Update, the statement “complete and circumferential membrane staining that is intense and within ≤10% of tumor cells” referred to an unusual pattern that did not need to be specified in the main portion of the figure. This information has now been moved to the footnote of Figure 1, which will now read as follows:

Unusual staining patterns of HER2 by IHC can be encountered that are not covered by these definitions. In practice, these patterns are rare and if encountered should be considered IHC 2+ equivocal. As one example, some specific subtypes of breast cancers can show IHC staining that is moderate to intense but incomplete (basolateral or lateral) and can be found to be HER2 amplified.22 Another example is circumferential membrane IHC staining that is intense but within ≤ 10% of tumor cells (heterogeneous but very limited in extent).

Consequently, the revised definition of IHC 2+ (HER2 equivocal) in this 2018 Focused Update (Figure 1) reflects a commonly accepted definition of invasive breast cancer that now reads “weak to moderate complete membrane staining observed in > 10% of tumor cells.” During the open comment period, pathologists requested guidance about the uncommon scenario of cases in which “intense circumferential membrane staining is observed in ≤ 10% of tumor cells.” As described in the footnote in Figure 1, such cases are relatively uncommon and are not typically subject to reflex testing.
Clinical Question 2

Must HER2 testing be repeated on a surgical specimen if initially negative test on core biopsy?

2013 Recommendation.—In Table 2 in the 2013 HER2 Testing Update, it was stated that, on the basis of some criteria (including a tumor grade 3), “If the initial HER2 test result in a core needle biopsy specimen of a primary breast cancer is negative, a new HER2 test must be ordered on the excision specimen…”

Revised 2018 Recommendation.—In the revised Table 2, it is now stated that, on the basis of some criteria (including a tumor grade 3), “If the initial HER2 test result in a core needle biopsy specimen of a primary breast cancer is negative, a new HER2 test may be ordered on the excision specimen…”

Clinical Question 3

Should invasive cancers with an HER2/CEP17 ratio of ≥2.0 but an average HER2 copy number of <4.0 per cell be considered ISH positive?

2013 Recommendation.—Cases in which the HER2/CEP17 ratio is ≥2.0 with an average HER2 signals per cell of <4.0 were considered ISH positive.

Revised 2018 Recommendation.—If a case has an HER2/CEP17 ratio of ≥2.0 but the average HER2 signals per cell is <4.0, a definitive diagnosis will be rendered based on additional workup.

If not already assessed by the institution or laboratory performing the ISH test, IHC testing for HER2 should be performed using sections from the same tissue sample used for ISH, and the slides from both ISH and IHC should be reviewed together to guide the selection of areas to score by ISH (local practice considerations will dictate the best procedure to accomplish this concomitant review):

- If the IHC result is 3+, diagnosis is HER2 positive.
- If the IHC result is 2+, recount ISH by having an additional observer, blinded to previous ISH results, count at least 20 cells that include the area of invasive cancer with IHC 2+ staining:
  - If reviewing the count by the additional observer changes the result into another ISH category, the result should be adjudicated per internal procedures to define the final category.
  - If the count remains an average of <4.0 HER2 signals per cell and the HER2/CEP17 ratio is ≥2.0, diagnosis is HER2 negative with a comment.
- If the IHC result is 0 or 1+, diagnosis is HER2 negative with a comment.

The Expert Panel recommends the following comment: evidence is limited on the efficacy of HER2-targeted therapy in the small subset of cases with an HER2/CEP17 ratio of ≥2.0 and an average HER2 copy number of <4.0 per cell. In the first generation of adjuvant trastuzumab trials, patients in this subgroup who were randomly assigned to the trastuzumab arm did not seem to derive an improvement in disease-free or overall survival, but there were too few such cases to draw definitive conclusions. IHC expression for HER2 should be used to complement ISH and define HER2 status. If the IHC result is not 3+ positive, it is recommended that the specimen be considered HER2 negative because of the low HER2 copy number by ISH and the lack of protein overexpression (Type: evidence-based; evidence quality: intermediate; strength of recommendation: strong). An algorithm for Clinical Question 3 is presented in Figures 3 and 4.

Literature Review and Analysis.—Members of the HER2 testing Expert Panel in 2013 had expressed concern about describing an invasive breast cancer as HER2 positive on the basis of a single HER2 ISH test that showed an HER2/CEP17 ratio of ≥2.0 but an average HER2 copy number of <4.0 signals per cell (Figure 3, group 2) and recommended additional testing of such cases. Members of the 2013 Guideline Update Panel also expressed their view that using the HER2/CEP17 ratio alone could be misleading in cases with CEP17 gains or losses and could lead to an underestimation or overestimation of HER2 amplification. However, the eligibility criteria for the first adjuvant trials of trastuzumab generally followed US Food and Drug Administration criteria (IHC 3+ or ISH ratio ≥2.0) regardless of...
average HER2 copy number based on HER2 signals per cell), and the panel in 2013 ultimately opted to consider these rare group 2 patients as having HER2-positive disease.

Since then, investigators have further reported on the outcome of patients with a group 2 dual-probe ISH test result. Greater experience and a more refined collection of test results in the past few years confirmed that such cases are infrequent (Table 3) and represented a small number of patients enrolled in the initial adjuvant trastuzumab trials. Among 4340 patients (41.5% of 10,468) screened by dual-probe fluorochrome ISH (FISH) for trials Breast Cancer International Research Group (BCIRG) 005 (HER2-negative trial) and BCIRG-006 (HER2-positive trial) and found to have a dual-probe ISH ratio of ≥2.0, only 71 (0.7% of 10,468) had an average number of HER2 signals per cell of <4.0. Furthermore, in 35 of these 71 patients who were also tested later by IHC, only 3 were IHC 2+ and none were IHC 3+. A retrospective assessment of potential benefit from trastuzumab in group 2 patients produced an observed hazard ratio estimate of slightly >1.0 (favoring no trastuzumab benefit), but the sample size was insufficient to statistically rule out a benefit from adjuvant trastuzumab in this group; nor could it be established statistically whether HER2-negative disease treated with just chemotherapy. In the HERA trial, outcomes different from patients with HER2-negative HER2-positive disease (type: evidence based; evidence quality: intermediate; strength of recommendation: strong). An algorithm for Clinical Question 4 is presented in Figures 3 and 5.

Clinical Question 4

Should invasive cancers with an average HER2 copy number of ≥6.0 signals per cell but an HER2/CEP17 ratio of <2.0 be considered ISH positive?

2013 Recommendation.—Cases in which the HER2/CEP17 ratio is <2.0 with an average of ≥6.0 HER2 signals per cell were considered ISH positive.

Revised 2018 Recommendation.—If a case has an average of ≥6.0 HER2 signals per cell with an HER2/CEP17 ratio of <2.0, formerly diagnosed as ISH positive for HER2, a definitive diagnosis will be rendered based on additional workup.

If not already assessed by the institution or laboratory performing the ISH test, IHC testing for HER2 should be performed using sections from the same tissue sample used for ISH, and the slides from both ISH and IHC should be reviewed together to guide the selection of areas to score by ISH (local practice considerations will dictate the best procedure to accomplish this concomitant review):

- If the IHC result is 3+, diagnosis is HER2 positive.
- If the IHC result is 2+, recount ISH by having an additional observer, blinded to previous ISH results, count at least 20 cells that include the area of invasion with IHC 2+ staining:
  - If reviewing the count by the additional observer changes the result into another ISH category, the result should be adjudicated per internal procedures to define the final category.
  - If the HER2/CEP17 ratio remains <2.0 with ≥6.0 HER2 signals per cell, diagnosis is HER2 positive.
- If the IHC result is 0 or 1+, diagnosis is HER2 negative with a comment.

The Expert Panel recommends the following comment: there are insufficient data on the efficacy of HER2-targeted therapy in cases with an HER2 ratio of <2.0 in the absence of protein overexpression because such patients were not eligible for the first generation of adjuvant trastuzumab clinical trials. When concurrent IHC results are negative (0 or 1+), it is recommended that the specimen be considered HER2 negative (type: evidence based; evidence quality: intermediate; strength of recommendation: strong). An algorithm for Clinical Question 4 is presented in Figures 3 and 5.

Literature Review and Analysis.—Based on available data, samples with ISH results in this category (ratio <2.0 and mean HER2 signals per cell ≥6.0) are uncommon, only representing between 0.4% and 3.0% of cases sent for dual-probe FISH testing (Table 3). These ISH cases have increases in both HER2 and control centromere signals, resulting in ratio results of <2.0. At the time of the pivotal HER2 trials, cases with these results were considered to have duplication of CEP17 (polysomy) and were most often excluded because they were considered negative for HER2 gene amplification, although the BCIRG-006 (Clinicaltrials.gov identifier: NCT00021255) allowed patients to be enrolled if the HER2 copy number was >10 and central IHC testing was 3+. Subsequent studies examining multiple regions of chromosome 17 supported that the majority of cases with these results have HER2 amplifications that include regions encompassing the centromere rather than true polysomy for the entire chromosome 17 (coamplification of control and HER2 signals). Based on these data, the 2013 Guideline Update clarified that cases with an average HER2 copy number of ≥6.0 HER2 signals per cell ISH results (by either single- or dual-probe assays) should be reported as HER2 positive by gene amplification. However, it was acknowledged that data on the clinical response of this group to HER2-targeted therapies were limited.

Since the 2013 update, additional data have been published including concurrent IHC results for this ISH category, and they show that this group can be heterogeneous. Data from a reanalysis of the HERA trial identified a highly amplified subgroup representing between 0.4% and 3.0% of cases sent for dual-probe FISH testing (Table 3). These ISH cases have increases in both HER2 and control centromere signals, resulting in ratio results of <2.0. At the time of the pivotal HER2 trials, cases with these results were considered to have duplication of CEP17 (polysomy) and were most often excluded because they were considered negative for HER2 gene amplification, although the BCIRG-006 (Clinicaltrials.gov identifier: NCT00021255) allowed patients to be enrolled if the HER2 copy number was >10 and central IHC testing was 3+. Subsequent studies examining multiple regions of chromosome 17 supported that the majority of cases with these results have HER2 amplifications that include regions encompassing the centromere rather than true polysomy for the entire chromosome 17 (coamplification of control and HER2 signals). Based on these data, the 2013 Guideline Update clarified that cases with an average HER2 copy number of ≥6.0 HER2 signals per cell ISH results (by either single- or dual-probe assays) should be reported as HER2 positive by gene amplification. However, it was acknowledged that data on the clinical response of this group to HER2-targeted therapies were limited.

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Since the 2013 update, additional data have been published including concurrent IHC results for this ISH category, and they show that this group can be heterogeneous. Data from a reanalysis of the HERA trial identified a small number of cases (21 total) originally considered negative due to ratios of <2.0 but with an average of ≥6.0 HER2 signals per cell. All of these cases had >3 mean CEP17 signals per cell, and 75% of them (15 of 20) had HER2 overexpression by IHC.

In a combined study of three major academic medical centers performing HER2 FISH and IHC, similar results were seen with 63 cases in this ISH category; 31.7% were IHC 3+ for HER2 by IHC, 55% were IHC 2+, and 13.7% were IHC 0 or 1+. This study also reported a higher frequency of Nottingham grade 3 cancers with these ISH results than with other ISH result categories. Published data from a reference laboratory at the University of Southern California described 48 cases with the same ISH characteristics and found that only 8.3% were IHC 3+, while 14.6% were IHC 2+ and 77% were IHC 0 or 1+. Additional analysis of these cases identified a highly amplified subgroup (eight total cases) with an average of 12.3 HER2
signals per cell that correlated well with HER2 IHC 2+ or 3+ (75%). This subgroup differed significantly from the other subgroup (40 total cases) that had a lower average of 6.8 HER2 signals per cell and 87.5% IHC negative (0 or 1+) results. Similarly, in the BCIRG central testing clinical trial data, of the limited cases (nine total) with IHC data and ISH results in this category, one was IHC 3+ positive, one was IHC 2+, and seven were IHC negative.10 Taken together, these results suggest that cases in this ISH category form a heterogeneous group that is best discriminated by the combination of IHC and ISH.

Due to the rarity of cases with these ISH results, there is still limited clinical evidence regarding benefit from HER2-targeted therapy. The BCIRG-005 data (no HER2-targeted treatment) indicated a worse disease-free and overall survival for this ISH category than for cases with both an HER2/CEP17 ratio of <2.0 and <4.0 signals per cell (ISH nonamplified).10 However, the few cases enrolled in the BCIRG-006 adjuvant trastuzumab trial with these ISH results were insufficient to assess whether there was benefit from HER2-targeted therapy, and statistical analysis was not attempted.

Overall, the absence of robust clinical data to guide decisions, and the variability in IHC data, support the concept that protein expression results should be used concurrently in this setting to aid in determining the significance of ISH results. In summary, group 3 cases are uncommon and heterogeneous. Based on available data, the ratio may not be a reliable indicator of the true gene amplification status.29–31 Given the evidence that some group 3 cases have true HER2 amplification rather than polyploidy for chromosome 17, particularly when the HER2 copy number is high, the Expert Panel ultimately favored continuing to classify these cases as HER2 positive unless the concurrent IHC result is clearly negative (0 or 1+).25–31

Repeat testing of other tissue samples from the patient may also be appropriate in this setting, and, in particularly challenging cases or if the results are in question, expert consultation may be appropriate and include alternative probes or other genetic methods.13 However, alternative probes should not be used as standard practice in view of the absence of outcome data.

**Clinical Question 5**

What is the appropriate diagnostic workup for invasive cancers with an average HER2 copy number of ≥4.0 but <6.0 signals per cell and an HER2/CEP17 ratio of <2.0, and initially deemed to have an equivocal HER2 ISH test result?

**2013 Recommendation.**—Cases in which the HER2/CEP17 ratio is <2.0 with an average HER2 copy number of ≥4.0 and <6.0 signals per cell were considered ISH equivocal, and additional workup was required (“Must order a reflex test [same specimen using IHC], test with alternative ISH chromosome 17 probe, or order a new test [new specimen if available, IISH or IHC]”).

**Revised 2018 Recommendation.**—If the case has an average HER2 signals per tumor cell of ≥4.0 and <6.0 and the HER2/CEP17 ratio is <2.0, formerly diagnosed as ISH equivocal for HER2, a definitive diagnosis will be rendered based on additional workup.

If not already assessed by the institution or laboratory performing the ISH test, IHC testing for HER2 should be performed using sections from the same tissue sample used for ISH, and the slides from both ISH and IHC should be reviewed together to guide the selection of areas to score by ISH (local practice considerations will dictate the best procedure to accomplish this concomitant review):

- If the IHC result is 3+, diagnosis is HER2 positive.
- If the IHC result is 2+, recount ISH by having an additional observer, blinded to previous ISH results, count at least 20 cells that include the area of invasion with IHC 2+ staining:
  - o If reviewing the count by the additional observer changes the result into another ISH category, the result should be adjudicated per internal procedures to define the final category.
  - o If the count remains an average of ≥4.0 and <6.0 HER2 signals per cell with an HER2/CEP17 ratio of <2.0, diagnosis is HER2 negative with a comment.
- If the IHC result is 0 or 1+, diagnosis is HER2 negative with a comment.

The Expert Panel recommends the following comment: It is uncertain whether patients with an average of ≥4.0 and <6.0 HER2 signals per cell and an HER2/CEP17 ratio of <2.0 benefit from HER2-targeted therapy in the absence of protein overexpression (IHC 3+). If the specimen test result is close to the ISH ratio threshold for positive, there is a high likelihood that repeat testing will result in different results by chance alone. Therefore, when IHC results are not 3+ positive, it is recommended that the sample be considered HER2 negative without additional testing on the same specimen (Type: evidence based; evidence quality: intermediate; strength of recommendation: strong). An algorithm for Clinical Question 5 is presented in Figures 3 and 6.

**Literature Review and Discussion.**—Cases with an average of ≥4.0 and <6.0 HER2 signals per cell and an HER2/CEP17 ratio of <2.0 were considered equivocal in the 2013 Guideline Update. This category (group 4 cases) has been reconsidered by the Expert Panel based on published literature since then, and was discussed by the representatives of expert laboratories and Expert Panel members during the open portion of the November 2016 in-person meeting. In many published studies, the incidence of equivocal cases has changed since the 2013 update, when more stringent requirements for ISH interpretation were described.14 The number of such cases within a laboratory varies based on the patient population referred for ISH testing, but it seems to be approximately 5% of cases (range, 1% to 16%).15–19,23–32 The use of alternative probes to adjudicate these cases has also increased since 2013.

Data from a central reference laboratory at Mayo Clinic included FISH data in a population of patients that is enriched from those with HER2 IHC 2+ results based on the original IHC testing performed locally by the referring laboratories (1922 patients; 85% IHC 2+). Among these cases tested by FISH at Mayo, 14% of patients had ISH equivocal results and one half became HER2 positive by ratio when a locally developed and analytically validated 17p arm probe (D17S122; Mayo Clinic, Rochester, MN) was combined with the HER2 probe (Abbott Molecular, Abbott Park, IL) for additional FISH testing. However, clinical information about benefit from HER2-targeted therapy in such patients is not available and may not exist because these patients would not have been eligible for the original pivotal trials.11 The reference laboratory experience reported by Press et al22 involving a different patient population found 4.6% of patients among 7,526 cases with equivocal
results when using the 2013 criteria, while 89% of these cases were IHC HER2 0 or 1+, 10% were IHC HER2 2+, and only 0.9% were IHC HER2 3+ positive. Another academic laboratory experience that combined results from three laboratories had a similar frequency of numbers of specimens with equivocal results (5.2%) among 8068 patients. Similar clinical characteristics were observed in patients with an average of ≥4.0 and <6.0 HER2 signals per cell, regardless of whether the ratio was above or below 2.0, and most of these cases were HER2 negative by IHC and more likely to be estrogen receptor positive.16

Group 4 cases reported as HER2 equivocal since the 2013 Guideline Update have posed a challenge to oncologists and patients due to a perceived ambivalence about whether to recommend HER2-targeted therapy. In the absence of an unequivocally positive or negative test result, multiple testing of the same tissue sample has been performed frequently, and many laboratories have relied exclusively on alternative probe testing to resolve cases that are more difficult. This has often included ISH testing using multiple chromosome 17 probes at once, many not analytically or clinically validated. Such indiscriminate testing often results in four or more ISH ratios being described in a single test report and a final designation of HER2 gene amplified if just a single ratio is ≥2.0. After careful consideration of this practice and available data, the 2018 Expert Panel strongly recommends against this as a routine testing strategy. When the HER2 ratio score is near a decision threshold (positive or negative), based on random variation in scoring, a subsequent test may result in a positive or negative score barely crossing the threshold (on either side). In such cases, repeated ISH testing may therefore not result in higher confidence in the final result.

Clinical correlation with other factors in a particular case (such as grade and special histologic subtypes) or repeat testing of other tissue samples from the patient may also be appropriate in this setting. In particularly challenging cases or if the results are in question, expert consultation may be appropriate and may include alternative probes or other genetic methods.11 However, alternative probes should not be used as standard practice due to limited data on outcomes for this subset of patients.33

**ADDITIONAL RESOURCES**

Additional information, including data supplements, evidence tables, and clinical tools and resources, can be found at www.asco.org/breast-cancer-guidelines. Patient information is available there and at www.cancer.net.

**Related ASCO Guidelines**


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**ASCO/CAP HER2 Testing in Breast Cancer Update—Wolff et al 1379**

Arch Pathol Lab Med—Vol 142, November 2018
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guidelines in breast cancer-re-evaluation of HERA trial fluorescence in situ


2013 American Society of Clinical Oncology guideline adaptation of the Cancer Care


AUTHORS’ DISCLOSURES OF POTENTIAL CONFLICTS OF INTEREST


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