An International Ki67 Reproducibility Study


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Background

In breast cancer, immunohistochemical assessment of proliferation using the marker Ki67 has potential use in both research and clinical management. However, lack of consistency across laboratories has limited Ki67's value. A working group was assembled to devise a strategy to harmonize Ki67 analysis and increase scoring concordance. Toward that goal, we conducted a Ki67 reproducibility study.

Methods

Eight laboratories received 100 breast cancer cases arranged into 1-mm core tissue microarrays—one set stained by the participating laboratory and one set stained by the central laboratory, both using antibody MIB-1. Each laboratory scored Ki67 as percentage of positively stained invasive tumor cells using its own method. Six laboratories repeated scoring of 50 locally stained cases on 3 different days. Sources of variation were analyzed using random effects models with log2-transformed measurements. Reproducibility was quantified by intraclass correlation coefficient (ICC), and the approximate two-sided 95% confidence intervals (CIs) for the true intraclass correlation coefficients in these experiments were provided.

Results

Intralaboratory reproducibility was high (ICC = 0.94; 95% CI = 0.93 to 0.97). Interlaboratory reproducibility was only moderate (central staining: ICC = 0.71, 95% CI = 0.47 to 0.78; local staining: ICC = 0.59, 95% CI = 0.37 to 0.68). Geometric mean of Ki67 values for each laboratory across the 100 cases ranged 7.1% to 23.9% with central staining and 6.1% to 30.1% with local staining. Factors contributing to interlaboratory discordance included tumor region selection, counting method, and subjective assessment of staining positivity. Formal counting methods gave more consistent results than visual estimation.

Conclusions

Substantial variability in Ki67 scoring was observed among some of the world’s most experienced laboratories. Ki67 values and cutoffs for clinical decision-making cannot be transferred between laboratories without standardizing scoring methodology because analytical validity is limited.


Uncontrolled proliferation is a key feature of malignancy. The nuclear proliferation marker Ki67 is of interest for various potential uses in the clinical management of breast cancer (eg, prognosis, prediction, and monitoring of response) (1–9). The most commonly used assay to assess Ki67 is immunohistochemical (IHC) staining with the MIB-1 antibody. However, interlaboratory methodology is inconsistent, and, despite the apparent prognostic utility of Ki67, routine use of this tumor biomarker has not been widely recommended by consensus guidelines panels such as that convened by the American Society of Clinical Oncology, mainly because of concerns regarding analytical validity (10).

With the goal of harmonizing Ki67 analytical methodology, Dowsett et al., on behalf of the International Ki67 in Breast Cancer Working Group of the Breast International Group and North American Breast Cancer Group, provided an overview of the current state of the art of Ki67 evaluation and proposed a set of guidelines for analysis and reporting of Ki67 (1). Although those guidelines aimed to reduce preanalytical and analytical variations, the Working Group recognized that actual scoring procedures varied substantially, contributing to a lack of consensus regarding optimal cutoffs that should be applied in various research and clinical decision-making settings. This lack of consistency has prevented direct comparisons of Ki67 across laboratories and clinical trials.

In an effort to harmonize Ki67 analysis, the Working Group studied intra- and interlaboratory reproducibility of IHC assays for Ki67 in breast cancer among a group of highly experienced pathology laboratories. A secondary aim was to identify key sources of variation, particularly those introduced by different scoring methodologies.
Methods

One hundred breast cancer cases were arranged into 1-mm single core tissue microarrays (TMAs), with 50 cases represented on each of two TMA blocks. Specimens were representative clinical cases of invasive breast carcinomas diagnosed in British Columbia during 2009 and 2010, 50 from an academic teaching hospital and 50 from a community hospital. Cases were centrally reviewed; 79% were estrogen receptor positive, and Nottingham grade (11) distribution was 32% grade 1, 44% grade 2, and 24% grade 3. The study was approved by the BC Cancer Agency Clinical Research Ethics Board (protocol H10-03420). Cases were anonymized (treatment, outcome, or follow-up are not part of this study), and the requirement for informed consent was waived.

Eight laboratories from North America and Europe participated. Each laboratory director has a track record of publishing one or more peer-reviewed papers regarding the clinical utility of Ki67. Three methods were conducted: one examining intralaboratory variability (Experiment 1) and two examining interlaboratory variability (Experiment 2, parts A and B).

Six laboratories participated in the intralaboratory reproducibility experiment (Experiment 1). Each laboratory used its own local protocol to stain one section from a 50-case TMA block, and then the laboratory scored Ki67 on this slide, using its own standard scoring method, on 3 separate days.

Eight laboratories participated in the interlaboratory reproducibility experiments (Experiment 2). Each laboratory received two sets of TMA sections, each set containing the same 100 cores: one centrally stained for Ki67 (Experiment 2A), the second stained by each laboratory following its own local protocol within 2 weeks of cutting (Experiment 2B).

Thus, Experiment 2A assessed intralaboratory reproducibility of Ki67 on centrally stained slides, eliminating variability of staining method (although this experiment was still subject to biological heterogeneity between serial sections cut from the same TMA). Experiment 2B assessed interlaboratory reproducibility of Ki67 when both local staining and local scoring methods were used (locally stained sections from Experiment 1 formed a subset of those included in Experiment 2B).

Details of each lab’s staining methodology are provided in Table 1. Central staining used the MIB-1 clone (mouse monoclonal antibody; Dako, Carpinteria, California). All labs also used MIB-1 from Dako for local staining, but dilution, incubation, and antigen retrieval and detection systems varied, as did hematoxylin counterstain supplier and time.

Ki67 was scored as the percentage of invasive tumor cells positively stained.

Statistical Analysis

Ki67 data were visualized using boxplots and dot plots. Pairwise intralaboratory and interlaboratory concordance were plotted using Bland–Altman plots, which graph the difference in Ki67 between any two paired observations against their mean (12). If there is high agreement in Ki67, the differences are expected to be centered about zero, with a small standard deviation. Crossed random effects models were fitted to formally quantify the amount of variability in Ki67 measurements contributed by each source of variation in the three experiments (13). Details are provided in the Supplementary Methods (available online). Because these random effects models rely on the data being normally distributed with constant variance, Ki67 data were log2-transformed to approximate a normal distribution and stabilize the variance (1). Specifically, a value of 0.1% was first added, and then a log base 2 transformation was applied to all observations. For example, for a Ki67 score of 30%, the recorded transformed value would be log2(30.1).

The intraclass correlation coefficient (ICC) was used as a summary measure of reproducibility. The ICC has a range of 0 to 1, with 1 denoting highest agreement. For Experiment 1, we estimated the ICC for repeated Ki67 measurements made on the same patient by the same laboratory (Supplementary Methods, available online). A credible interval of the ICC was obtained using Markov Chain Monte Carlo routines for fitting generalized linear mixed models, using the MCMCglmm package in R (14). For Experiments 2A and 2B, we estimated the ICC for Ki67 measurements made on the same patient by different laboratories (Supplementary Methods, available online). The approximate two-sided 95% confidence intervals (CIs) for the true ICCs in these experiments were computed using a closed-form formula (15). This ICC has a particularly intuitive interpretation as the proportion of the total variance in the measurements across patients and labs attributable to the true biological variability between patients.

All data analyses were performed using the R language version 2.11.1 (16). The crossed random effects models described in Model 1 and Model 2 (Supplementary Methods, available online) were fitted using the lme4 package (17).

Results

Intralaboratory Reproducibility (Experiment 1)

Figure 1 presents Bland–Altman plots for each possible pairing of Ki67 scores (on the original scale as percentage of positively stained tumor cells) measured on different days by the same laboratory. Four laboratories (Laboratories C, D, E, and H) exhibited the highest degree of internal consistency. Two of these laboratories (Laboratories E and H) used formal point (individual cell) counting methods, whereas Laboratories C and D used visual estimation. Laboratory C was clearly estimating in increments of 5%, at estimates of 5% or greater, and this rounding may have contributed to a heightened impression of internal consistency. For further analyses, we considered Laboratories D, E, and H to have best demonstrated internal consistency. Supplementary Table 1 (available online) provides summary statistics for log2-transformed Ki67 scores by laboratory and day.

The estimates of the variance terms (ν1, ν2, ν3, 0) in Model 1 (Supplementary Methods, available online) were 1.91, 0.29, 0.39, and 0.14, respectively. The largest source of the total variation appears to have been patient biology (as expected, reflecting biological differences in Ki67 levels among different tumors), followed by the interaction between patient and laboratory effects, laboratory, and residual errors. The ICC estimate was 0.94 ([1.91 + 0.29 + 0.39] / [1.91 + 0.29 + 0.39 + 0.14] = 0.94; 95% CI = 0.93 to 0.97). Therefore, the intralaboratory reproducibility was very high, indicating that the laboratories could deliver internally consistent results.
<table>
<thead>
<tr>
<th>Lab</th>
<th>Scoring method</th>
<th>KI67 staining (all used MIB-1 antibody, from Dako)</th>
<th>Counterstain</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Antigen retrieval</td>
<td>MIB-1 dilution</td>
</tr>
<tr>
<td>Lab A</td>
<td>Visual estimate 1% increments (&lt;10%), 5% increments (10%–30%), 10% increments (&gt;30%)</td>
<td>High pH in PT Link 1:200 20 min Dako Flex</td>
<td>Harris Hem Intermedico</td>
</tr>
<tr>
<td>Lab B</td>
<td>Visual estimate (1% increments)</td>
<td>Pressure cooker in citrate buffer pH 6 for 6 min 1:50 30 min Dako Envision</td>
<td>Harris acidic, Thermo Fisher</td>
</tr>
<tr>
<td>Lab C</td>
<td>Visual estimate 1% increments (&lt;5%), 5% increments (10%–40%), 10% increments (&gt;40%)</td>
<td>Ventana CC1 short (30 min) 1:100 32 min 37°C UltraView</td>
<td>Hematoxylin Ventana/Roche reference:760–2021</td>
</tr>
<tr>
<td>Lab D</td>
<td>Visual estimate (1% increments)</td>
<td>Ventana CC1 EDTA pH 8 1:50 32 min UltraMap Dako Envision</td>
<td>Hematoxylin Gill 1, Ventana Dako product No. S3301</td>
</tr>
<tr>
<td>Lab E</td>
<td>Counting 500 cells using a hemocytometer; any level of stain is positive</td>
<td>Leica BONDmax TRIS pH 9 Tris-EDTA 1:100 15 min Leica BONDmax DAB</td>
<td>Leica kit</td>
</tr>
<tr>
<td>Lab F</td>
<td>Counting ~200 cells using a keypad Web application</td>
<td>Leica BONDmax TRIS pH 9 Tris-EDTA 1:250 20 min UltraVision</td>
<td>Type: Mayer (75% Mayer, 25% water) Supplier: BBC Biocare product No.: 3580</td>
</tr>
<tr>
<td>Lab G</td>
<td>Visual estimate 10% increments (exceptions made for 1% and 95%; 28 cases scored in 1% increments and one case scored as 95%)</td>
<td>Microwave Dako Epitope Retrieval Solution 10 min 1:50 20 min Dako Real</td>
<td>Haemalum Mayer TCS Biosciences Ltd. catalog No. HS315</td>
</tr>
<tr>
<td>Lab H</td>
<td>Counting entire invasive tumor in each core using a 10×10 grid in a ×10 eyepiece graticule</td>
<td>Microwave Dako Epitope Retrieval Solution 10 min 1:50 20 min Dako Real</td>
<td>Haemalum Mayer TCS Biosciences Ltd. catalog No. HS315</td>
</tr>
</tbody>
</table>

* EDTA = Ethylenediaminetetraacetic acid; PT = PT Link is a pre-treatment system from Dako; TFS = Thermo Fisher Scientific; TRIS = Tris(hydroxymethyl)aminomethane.
Interlaboratory Reproducibility

Experiment 2A (Central Staining, Local Scoring Method).

Table 2 provides, for each laboratory, summary statistics of log2-transformed Ki67 scores observed when all laboratories used their own local scoring methods on centrally stained sections. The geometric mean of Ki67 (by taking the antilogs of the means on log2 scale) across the 100 cases ranged from 7.1% (Laboratory G) to 23.9% (Laboratories D and F) (the arithmetic mean ranged from 15.6% to 31.1%). Such a range indicates substantial differences in Ki67 measurement across laboratories on centrally stained slides from the same cases. Figure 2A presents side-by-side boxplots. Figure 3A plots the individual Ki67 measurements made for each of the 100 patients by each laboratory. Some laboratories tended to score Ki67 at several discrete values, whereas others tended to score on a more continuous scale (Figure 3A). In general, Figure 2 and Figure 3 suggest that high variability in Ki67 scoring was detected among the laboratories.

Using statistical Model 2 (Supplementary Methods, available online), we obtained variance estimates to describe biological variation between patients and variation between laboratories. The estimates of these variances (ν', ν'' , θ') in Model 2 were 1.78, 0.34, and 0.39, respectively. The largest variation appears to have come from individual patients, followed by residual errors and laboratory. The ICC estimate was 0.71 (1.78 / [1.78 + 0.34 + 0.39] = 0.71;
95% CI = 0.47 to 0.78). Therefore, Ki67 achieved only moderate reproducibility across the laboratories when they used their own scoring methodology on sections stained in a central laboratory.

**Experiment 2B (Local Staining, Local Scoring Method).** Table 3 provides, for each laboratory, the summary statistics of log2-transformed Ki67 scores that were observed when all laboratories used their own local scoring methods on sections they stained according to their own local staining methods. The geometric mean of Ki67 across the 100 cases ranged from 6.1% (Laboratory G) to 30.1% (Laboratory F) (the arithmetic mean ranged from 12.6% to 35.5%), suggesting a large interlaboratory variability in Ki67 measurement.

Figure 2B presents side-by-side boxplots. Figure 3B presents a dot plot. In both figures, laboratories are ordered by increasing median Ki67 values. Again, there appears to be a large interlaboratory variability. Although laboratory order is not identical in Experiments 2A and 2B when laboratories are arranged by increasing median Ki67 values, the general distribution of Ki67 data for individual laboratories seems to be largely preserved between the two experiments (Figure 3).

To quantify the sources of variation in the data, we again fit a two-way crossed random effects model specified in Model 2 (Supplementary Methods, available online). The estimates of the variance terms (\(\nu_1^2, \nu_2^2, \theta^2\)) in Model 2 were 1.71, 0.46, and 0.71, respectively. Similar to Experiment 2A, the largest variation was attributed to biological variability between patients, followed by residual errors and laboratory-to-laboratory variability. Using the variance estimates, the ICC was computed as 0.59 (1.71 / [1.71 + 0.46 + 0.71] = 0.59; 95% CI = 0.37 to 0.68). Therefore, allowing each laboratory to stain the slides based on its own local staining protocol (in addition to scoring them based on its own local scoring method) further decreased the interlaboratory reproducibility compared with Experiment 2A.

The three laboratories that demonstrated the highest level of intralaboratory reproducibility in Experiment 1 (Laboratories D, E, and H) also tended to generate Ki67 values most similar to one another (Figure 4). In both experiments, Laboratory D (which used visual estimation) scored slightly higher in terms of median Ki67 index than Laboratory H, followed by Laboratory E (the latter two used formal counting methods).

In an exploratory analysis, we found that interlaboratory variation was lower among laboratories using formal counting approaches (\(n = 3\) labs) compared with those using visual estimation (\(n = 5\) labs). Spaghetti plots graphically demonstrate this lab-to-lab variability per case (Supplementary Figures 1, A and B, and 2, A and B, available online). ICC was also higher among labs using formal counting: 0.82 (95% CI = 0.19 to 0.87) by counting vs 0.72 (95% CI = 0.32 to 0.80) by visual estimation in Experiment 2A, and 0.75 (95% CI = 0.10 to 0.84) by counting vs 0.64 (95% CI = 0.31 to 0.72) by visual estimation in Experiment 2B. ICC confidence intervals are wide in these subgroup analyses because of the limited number of laboratories.

**Discussion**

In this study, we observed large variation among a group of highly experienced analysts in determination of levels of Ki67, a biomarker that has been incorporated into clinical care by pathology laboratories worldwide. Although intralaboratory reproducibility of Ki67 evaluation by experienced laboratories was generally good, suggesting that analytical validity may be achievable, interlaboratory reproducibility was only moderate and was even worse when both staining and scoring were done locally. These results support
Figure 3. Dot plot of Ki67 measurements by laboratory from Experiments 2A and 2B. A) Centrally stained, local scoring method. B) Locally stained, local scoring method. Laboratories are reordered according to increasing median Ki67 value (percentage of positively stained tumor cells). When multiple observations have the same Ki67 value, random jittering is used to displace the points vertically to aid visualization.
Ki67 levels might be used to determine prognosis or residual risk after primary therapy, to predict activity of systemic therapies, or to monitor patients for sustained response or resistance to delivered therapies (1). However, according to terminology suggested by the Evaluation of Genomic Applications in Practice and Prevention initiative (19), an assay cannot have clinical utility for any of these uses unless its analytical validity has been demonstrated. Our results suggest that even among some of the world’s experts in IHC staining and evaluation of Ki67, the analytical validity for this assay is unacceptably poor. Unless an individual pathology laboratory has demonstrated that its staining and scoring methodology, including cutoff determination, meet the highest level of evidence for clinical utility (20), clinicians should use Ki67 results with great caution.

Although interlaboratory differences in staining methodology contributed to Ki67 variability, we also observed a striking heterogeneity in scoring interpretation of centrally stained slides, even using a TMA platform that reduces concern about selection of tissue areas for reading. We did not assess Ki67 in core biopsies or whole sections, diagnostic formats that add complexities regarding which area to score and how to handle “hot spots” of proliferation. If high levels of interobserver concordance cannot be achieved with TMAs, it is even less likely they would be achievable using standard clinical diagnostic formats. Indeed, in a recently reported study in which breast cancer whole sections were distributed to 15 pathologists, Varga et al. observed problematically high interobserver variability among cases in the Ki67 midrange (Ki67 index of 8% to 15%), precisely the region in which most cutoffs are located for making clinical decisions (21). In their study, no single factor (counting method, threshold for positivity, area chosen to score, or staining methodology) explained these differences. There is strong evidence, however, that future approaches using agreed-upon consensus guidelines may improve observer variability and assist standardization (22), an issue that we will address in the next phase of our studies.

Even within the TMA format in our study, several other sources of variability contributed to poor interlaboratory agreement, including whether the laboratory used formal counting of nuclei vs visual estimation, unavoidably subjective assessments of which nuclei represent invasive cancer cells, and what threshold to use for “positive” staining. Although the contributions from these factors are not rigorously separable in our data, the data distributions do suggest that laboratories using formal counting methods gave more consistent results than those using visual estimation.

Clinical decision-making regarding treatment options in breast cancer often relies on the application of a Ki67 cutoff to classify patients into “Ki67 high” or “Ki67 low” risk groups. Widely varying cutoff values, however, further impede the clinical utility of Ki67 and make it difficult to compare Ki67 data across different studies. Reviews of multiple studies in early breast cancer show that cutoffs ranging from 0% to 28.6% have been used (23,24). The 2011 St. Gallen International Consensus Meeting Conference Panel recommended a cutoff of 13.5% to distinguish between “luminal A” and “luminal B/HER2-negative” subtypes in patients with node-negative invasive breast cancer (2). Our data suggest that even if a common Ki67 cutoff is agreed upon, lack of interlaboratory reproducibility in Ki67 measurements represents a major obstacle to confident use of Ki67 for clinical decisions. For example, if the cutoff of 13.5% were applied to the two laboratories that had substantially discordant Ki67 measurements from our Experiment 2A (central staining, local scoring method), 31 of 96 patients (32.3%) would be classified as “Ki67 high” by Laboratory D but as “Ki67 low” by Laboratory B (Figure 5). Further, when the 13.5% cutoff is applied to all Experiment 2A laboratories, the laboratory-specific percentage of patients who would be classified as luminal A varies widely: 56.0% (Laboratory A), 47.9% (Laboratory B), 74.0% (Laboratory C), 77.8% (Laboratory D), 57.0% (Laboratory E), 81.0% (Laboratory F), 30.3% (Laboratory G), 69.0% (Laboratory H). Application of this or other cutoffs for selecting patients for chemotherapy is inappropriate without rigorous analytical standardization.

Computerized digital image analysis has been suggested as a potential solution to problems of analytical subjectivity and interobserver variability in Ki67 assessment (25–28). We have intentionally limited our assessments to visual methods requiring no special equipment because these could be readily and inexpensively adopted by laboratories around the world. Image analysis methods could be a subject of future studies if visual assessments cannot achieve sufficient analytical validity. Recent studies in the neuroendocrine tumor literature, where Ki67 scoring is part of World Health Organization–recommended grading systems, report that digital image analysis (of this comparatively homogeneous tumor type) performs as well as or better than visual counting, with both superior to visual estimation (29–31).

### Table 3. Summary statistics of log2-transformed Ki67 measurements by laboratory, Experiment 2B (local staining, local scoring method)*

<table>
<thead>
<tr>
<th>Laboratory</th>
<th>Min</th>
<th>Q1</th>
<th>Median</th>
<th>Mean</th>
<th>Q3</th>
<th>Max</th>
<th>SD</th>
<th>No. of missing observations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laboratory A</td>
<td>0.14</td>
<td>3.34</td>
<td>3.92</td>
<td>3.74</td>
<td>4.33</td>
<td>6.65</td>
<td>1.42</td>
<td>1</td>
</tr>
<tr>
<td>Laboratory B</td>
<td>−3.32</td>
<td>1.63</td>
<td>3.92</td>
<td>3.62</td>
<td>5.65</td>
<td>6.63</td>
<td>2.29</td>
<td>9</td>
</tr>
<tr>
<td>Laboratory C</td>
<td>−3.32</td>
<td>2.35</td>
<td>3.92</td>
<td>3.54</td>
<td>4.85</td>
<td>6.65</td>
<td>2.08</td>
<td>2</td>
</tr>
<tr>
<td>Laboratory D</td>
<td>1.63</td>
<td>3.92</td>
<td>4.81</td>
<td>4.56</td>
<td>5.33</td>
<td>6.62</td>
<td>1.17</td>
<td>1</td>
</tr>
<tr>
<td>Laboratory E</td>
<td>1.63</td>
<td>3.34</td>
<td>3.96</td>
<td>3.94</td>
<td>4.15</td>
<td>6.49</td>
<td>1.01</td>
<td>2</td>
</tr>
<tr>
<td>Laboratory F</td>
<td>2.04</td>
<td>4.47</td>
<td>5.05</td>
<td>4.91</td>
<td>5.51</td>
<td>6.48</td>
<td>0.92</td>
<td>1</td>
</tr>
<tr>
<td>Laboratory G</td>
<td>0.14</td>
<td>0.14</td>
<td>2.35</td>
<td>2.6</td>
<td>4.33</td>
<td>6.49</td>
<td>1.85</td>
<td>2</td>
</tr>
<tr>
<td>Laboratory H</td>
<td>1.07</td>
<td>3.6</td>
<td>4.53</td>
<td>4.4</td>
<td>5.36</td>
<td>6.59</td>
<td>1.25</td>
<td>1</td>
</tr>
</tbody>
</table>

* max = maximum; min = minimum; Q1 = first quartile; Q3 = third quartile; SD = standard deviation.
The results from our study emphasize the major differences that exist even among experienced laboratories in results from Ki67 analyses in breast cancer and focus attention on the need for harmonization of scoring procedures if this biomarker’s potential is to be realized in breast cancer. We are currently studying whether a prespecified target of success (ICC = 0.9) can be achieved when scorers “train” using a calibration tool designed to mitigate causes of systematic Ki67 scoring differences. Because of widespread use of Ki67 in many research and clinical settings, however, we feel that it is imperative at the present time to report the high interlaboratory variability in Ki67 we observed in this study.

Our study is limited in the sense that it pertains specifically to analytical validity and does not touch on clinical validity. As stated above, we used TMA slides rather than specimen formats typically used in clinical practice (core biopsies or whole sections). Because we used different serial sections of the (same) TMA, some variability in scoring could conceivably be attributable to the section received (although we believe this to be minimal). Although lower than the variability introduced by scoring differences, variability introduced by methodological differences in the IHC staining processes could not be assigned to individual steps within this multistage procedure. Although we succeeded in underscoring an existing problem, we are not yet able to offer a solution. As we note above, we are actively carrying out follow-on studies examining the effect of training labs on a common scoring method, with a view to developing a standardized approach. If these studies are successful, we would extend our approach to core biopsies and whole sections and link findings to patient outcomes to confirm clinical utility.

In summary, although there are multiple potential applications for Ki67 in research and clinical management that are supported by an extensive literature (3,4,23), the clinical utility of Ki67 in breast cancer remains elusive because of analytical concerns. Variability among laboratories in their approaches to scoring is
a major contributor to discordance in results. We recommend that caution be exercised at present when comparing Ki67 results across different laboratories or studies, and we echo the sentiments of the 2007 American Society of Clinical Oncology Tumor Marker Guidelines Committee against using Ki67 in routine clinical practice (10). The following recommendation from our 2011 policy paper therefore still holds for assessment of Ki67 index in breast cancer, when performed by visual assessment of glass slides: “Cut points for prognosis, prediction, and monitoring should only be applied if the results from local practice have been validated against those in studies that have defined the cutoff for the intended use of the Ki67 result” (1).

References


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**Notes**

The study sponsors had no role in the design of the study; the collection, analysis, and interpretation of the data; the writing of the manuscript; and the decision to submit the manuscript for publication.

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