Recommendations for Validating Whole Slide Imaging Systems for Diagnostic Purposes in Pathology

Anil V Parwani, MD, PhD
On behalf of the CAP WSI Validation Expert Panel
Disclosure Information

- No disclosures.

- This presentation will include a discussion of devices that have not yet been approved by the FDA.
Digital Pathology Uses


- Primary diagnosis
- 2nd opinion (consultation)
- Telepathology
- Quality Assurance (PT)
- Archiving & Sharing
- Education/Conferencing
- Image analysis
- Research & Publications
- Marketing & Business
- Track (audit) & Training
A Better Lens on DISEASE

Computerized pathology slides may help doctors make faster and more accurate diagnoses.

BY MIKE MAY

Imaging Process

1. Capture
2. Save
3. Edit
4. Share
Imaging Modes

Static

Video

Robotic

WSI
Whole Slide Imaging

- WSI = Virtual microscopy (digital slides)
- Digitization of glass slides simulates light microscopy
- Provides access to all areas of interest on a slide
- High resolution digital images
- Digitization at multiple magnifications
- Scanning in multiple focal planes (x, y & z axes)
WSI scanners

WSI Scanner Components

- Optical microscope
- Slide rack
WSI Scanning Strategies

A - Tile-based acquisition mode (grid pattern)
B - Line-scan acquisition mode (linear pattern)

WSI Regulatory Issues

- No specific CLIA regulations for Digital Pathology
- FDA have designated WSI systems to be class III (highest risk) medical devices
Recommendations for Validating Whole Slide Imaging Systems for Diagnostic Purposes in Pathology
WSI Validation Expert Panel
Convened June 2010

Liron Pantanowitz, MD, Chair
Bruce Beckwith, MD
Alexis Carter, MD
Lydia Contis, MD
Andrew Evans, MD, PhD
Walter Henricks, MD
Christopher Otis, MD
Anil Parwani, MD
John Sinard, MD
Avtar Lal, MD, PhD
Lisa Fatheree, SCT(ASCP) CAP Staff
What needs to be done to “validate” a whole slide digital imaging system for diagnostic purposes before it is placed in clinical service?

Panel addressed: The intended use, preparation types, number of cases, equipment, personnel, and process.
Systematic Review Results

• Literature search conducted:
  o 767 studies met the search term requirement
  o 27 underwent data extraction for evidence evaluation

• Panel met 8 times to develop draft recommendations

• Open comment period (July - Aug 2011):
  o 132 respondents; 531 comments
  o Evidence tables not completed at that time

• Panel met 10 additional times to review feedback, make revisions to recommendations, and assess strength of evidence supporting the 12 final recommendations
Quality Assessment and Grading of Evidence

- **Strength of evidence**: level of evidence, quantity, size of the effect, statistical precision and, quality assessment (risk of bias) of included studies.

- Also taken into account were the study components of consistency, clinical impact, generalizability, and applicability to WSI when determining the strength of evidence score.
## Definitions of Grading of Recommendations

### Definition of Australian National Health and Medical Research Council (NHMRC) to grades of recommendations

<table>
<thead>
<tr>
<th>Grade of recommendation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Body of evidence can be trusted to guide practice</td>
</tr>
<tr>
<td>B</td>
<td>Body of evidence can be trusted to guide practice in most situations</td>
</tr>
<tr>
<td>C</td>
<td>Body of evidence provides some support for recommendation(s) but care should be taken in its application</td>
</tr>
<tr>
<td>D</td>
<td>Body of evidence is weak and recommendation must be applied with caution</td>
</tr>
<tr>
<td>Guidance</td>
<td>Description</td>
</tr>
<tr>
<td>----------------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Recommendation</td>
<td>For moderate and highest level of evidence (Grade A/B) or where statements are unlikely to change based on further evidence. Note: Can also be in the negative, i.e., ‘Recommend Against’ or ‘Not Recommended’.</td>
</tr>
<tr>
<td>Suggestion</td>
<td>For inconclusive, conflicting and/or weak evidence (Grade C) or where statements most likely correct but could be better supported by additional data.</td>
</tr>
<tr>
<td>Expert Consensus</td>
<td>There is a gap, poor evidence (Grade D) or no evidence to support statement but necessary to address the topic. May be qualified with “requires future studies to be conducted”.</td>
</tr>
<tr>
<td>No recommendation</td>
<td>No statement generated for this key question / topic.</td>
</tr>
</tbody>
</table>
Guideline Statement #1

• All pathology laboratories considering the implementation of WSI technology for clinical diagnostic purposes should carry out their own validation studies.

Grade: Expert Opinion
**Validation is NOT Verification**

- **Validation** = The process of testing a procedure or instrument to assess its performance & determine whether it is acceptable
  - This is necessary to prove that it performs as expected & achieves the intended result
  - Did I do the right thing?

- **Lab developed tests (LDTs)** are not FDA-approved (off-label use), and require validation

- **Verification** = Checking the manufacturer’s claims for performance specifications before use
  - Verifying that a product/test can be replicated in the lab before patient testing (directed by the manufacturer’s user manual)
  - Typically performed in a clinical lab wanting to implement an FDA-approved instrument or method
  - Did I do the thing right?
• Variables between institutions can affect performance.
• Manufacturer device validation (i.e. verification) alone is insufficient.
• Simple guidelines provided for cytology screening devices (which were FDA approved) will not suffice.
• Comment feedback: 87% agreement
Guideline Statement #2

- Validation should be appropriate for and applicable to the intended clinical use and clinical setting of the application in which WSI will be employed. Validation of WSI systems should involve specimen preparation types relevant to intended use (e.g., formalin-fixed paraffin-embedded tissue, frozen tissue, IHC stains, cytology slides, hematology blood smears).

Grade: Recommendation, Level A
## Different Outcomes of WSI and Glass Slides with Different Types of Preparation

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Preparations for WSI and Glass Slides</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>H&amp;E</td>
</tr>
<tr>
<td>Accuracy of WSI or glass slides</td>
<td>WSI</td>
</tr>
<tr>
<td></td>
<td>95%</td>
</tr>
<tr>
<td>Concordance between WSI and glass slides</td>
<td>84%</td>
</tr>
<tr>
<td>Discordance between WSI and glass slides</td>
<td>16%</td>
</tr>
<tr>
<td>Concordance and minor discordance between WSI and glass slides</td>
<td>97%</td>
</tr>
</tbody>
</table>
• **Note:** If a new intended use for WSI is contemplated, and this new use differs materially from the previously validated use, a separate validation for the new use should be performed.

• **For example:** A validation study used to support the diagnostic use of digitized slides for routine surgical pathology may not necessarily apply to the use of frozen section digitized slides (e.g., with tissue folds, more pale staining, more mounting medium, etc.).

• **Comment feedback:** 81% agreement; Panel revised original statement for clarity
Guideline Statement #3

• The validation study should closely emulate the real-world clinical environment in which the technology will be used.

Grade: Recommendation, Level A
• **Goal of validation:**
  - Conducted in a manner that mimics how WSI will be used in the specific lab’s work environment.
  - Mimic how the system is to be used after “go live”.

• **For example:** If rapid digitization of glass slides is required for clinical use (e.g. frozen sections), then timely preparation & reading of WSI should be included in the validation process.

• **Comment feedback:** 91% agreement
Different Outcomes of Whole Slide Imaging (WSI) and Glass Slides with Emulation of Real-World Clinical Environment

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>WSI</th>
<th>Glass Slides</th>
</tr>
</thead>
<tbody>
<tr>
<td>Accuracy of WSI</td>
<td>89%</td>
<td>92%</td>
</tr>
<tr>
<td>Concordance between WSI and glass slides</td>
<td>86%</td>
<td></td>
</tr>
<tr>
<td>Discordance between WSI and glass slides</td>
<td>14%</td>
<td></td>
</tr>
<tr>
<td>Concordance and minor discordance between WSI and glass slides</td>
<td>98%</td>
<td></td>
</tr>
</tbody>
</table>
Guideline Statement #4

• The validation study should encompass the entire WSI system.

• Note: It is not necessary to validate separately each individual component (e.g., computer hardware, monitor, network, scanner) of the system nor the individual steps of the digital imaging process.

Grade: Recommendation, Level B
• WSI system is made up of different components: scanner, hardware, software, network & viewing monitor (+ pathologist).

• Parameters of each of component may impact digital image quality and therefore interpretation.

• Imaging process involves several steps including image acquisition, storage, sharing & viewing.

• Recommend the entire WSI system & imaging process be validated.

• All components are important & should not be separated, including technical system (“tool”) & observer (“pathologist).

**Comment feedback: 89% agreement**
Different Outcomes of Whole Slide Imaging (WSI) and Glass Slides with Entire WSI System

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>WSI</th>
<th>Glass Slides</th>
</tr>
</thead>
<tbody>
<tr>
<td>Accuracy of WSI</td>
<td>89%</td>
<td>92%</td>
</tr>
<tr>
<td>Concordance between WSI and glass slides</td>
<td>83%</td>
<td></td>
</tr>
<tr>
<td>Discordance between WSI and glass slides</td>
<td>17%</td>
<td></td>
</tr>
<tr>
<td>Concordance and minor discordance between WSI and glass slides</td>
<td>98%</td>
<td></td>
</tr>
</tbody>
</table>
Guideline Statement #5

- Revalidation is required whenever a significant change is made to any component of the WSI system.

Grade: Expert Opinion
• Significant changes to a WSI system may affect the interpretation of digital slides.
  o e.g. new scanner, major hardware or software upgrade

• For major changes the validation process should be repeated:
  o With these new changes incorporated in the WSI system
  o To demonstrate that it can still be employed for the intended use

• Minor changes can be managed through a facilities change management procedure.

Comment feedback: 85% agreement
Guideline Statement #6

- A pathologist(s) adequately trained to use the WSI system must be involved in the validation process.

Grade: Recommendation, Level B
• Validation process should include individual(s) who will actually be using the system to make diagnoses.

• Published validation studies: Average # evaluators = 8 individuals/study (range, 3 - 26 persons).

• Validation team may include other pathology staff (e.g. image technician, histotechnologist, PA), IT personnel and/or consultants.

• User training is important, but not part of validation. Training methods are outside of the scope of this document.

• Comment feedback: 91% agreement
### Different Outcomes of WSI and Glass Slides with Respect to Training of Pathologists

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Training [Mean +/- SD] or Percentage</th>
<th>No Training</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intra-observer agreement of WSI</td>
<td>0.93 ± 0.05</td>
<td>NR</td>
</tr>
<tr>
<td>Intra-observer agreement of glass slides</td>
<td>0.93 ± 0.03</td>
<td>NR</td>
</tr>
<tr>
<td>Intra-observer agreement between WSI and glass slides</td>
<td>NR</td>
<td>0.71</td>
</tr>
<tr>
<td>Inter-observer agreement of WSI</td>
<td>0.82 ± 0.01</td>
<td>0.53 ± 0.11</td>
</tr>
<tr>
<td>Inter-observer agreement of glass slides</td>
<td>0.85 ± 0.01</td>
<td>0.59 ± 0.06</td>
</tr>
<tr>
<td>Accuracy of WSI</td>
<td>95%</td>
<td>79%</td>
</tr>
<tr>
<td>Accuracy of glass slides</td>
<td>99%</td>
<td>81%</td>
</tr>
<tr>
<td>Concordance between WSI and glass slides</td>
<td>89%</td>
<td>84%</td>
</tr>
<tr>
<td>Discordance between WSI and glass slides</td>
<td>11%</td>
<td>16%</td>
</tr>
<tr>
<td>Concordance and minor discordance between WSI and glass slides</td>
<td>98%</td>
<td>98%</td>
</tr>
<tr>
<td>Interpretation time of WSI (Min)</td>
<td>4.9 ± 1.6</td>
<td>11.5 ± 2.5</td>
</tr>
</tbody>
</table>
OPTIONAL: How many cases do you think should be included in a validation study?

1. None (validation is not necessary)
2. Less than 10 cases
3. 60 cases
4. 100 cases
5. 500 cases
6. More than 500 cases
Guideline Statement #7

- The validation process should include a sample set of at least 60 cases for one application (e.g., H&E stained sections of fixed tissue, frozen sections, cytology, hematology) that reflects the spectrum and complexity of specimen types and diagnoses likely to be encountered during routine operation.

  - Note: The validation process should include another 20 cases for each additional application (e.g., immunohistochemistry, special stains).

  Grade: Recommendation, Level A
Validation of WSI systems should:
- Involve specimen preparation types relevant to intended use
- Not specific organ systems, diseases, microscopic changes or diagnoses

Important that an adequate sample size be used to allow pathologists to negotiate any technology learning curve.

Literature: Average 92 cases/study (range 10 to 633 cases).

Comment feedback: 73% - Panel made revisions with evidence (e.g. previously 100 cases)
## Different Outcomes of WSI and Glass Slides with Different Number of Cases

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Average Number of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>20 cases</td>
</tr>
<tr>
<td>Accuracy of WSI</td>
<td>72%</td>
</tr>
<tr>
<td>Accuracy of glass slides</td>
<td>77%</td>
</tr>
<tr>
<td>Concordance between WSI and glass slides</td>
<td>75%**</td>
</tr>
<tr>
<td>Discordance between WSI and glass slides</td>
<td>25%</td>
</tr>
<tr>
<td>Concordance and minor discordance between WSI and glass slides</td>
<td>95%</td>
</tr>
</tbody>
</table>

* P< .001 vs accuracy of 200 cases glass slides

** P=.002 vs concordance of 60 cases & P<.001 vs concordance of 200 cases
Guideline Statement #8

- The validation study should establish diagnostic concordance between digital and glass slides for the same observer (i.e., intraobserver variability).

**Grade:** Suggestion, Level A

Due to the conflicting nature of the good quality evidence for agreement and concordance, the statement stands as a Suggestion.
Different Outcomes of WSI and Glass Slides with Intraobserver and Interobserver Agreement

<table>
<thead>
<tr>
<th>Outcome</th>
<th>WSI</th>
<th>Glass Slides</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intra-observer agreement of WSI or glass slides with reference standard</td>
<td>0.93 ± 0.05</td>
<td>0.93 ± 0.03</td>
</tr>
<tr>
<td>Intra-observer agreement of WSI and glass slides</td>
<td>0.71</td>
<td></td>
</tr>
<tr>
<td>Inter-observer agreement of WSI or glass slides</td>
<td>0.68 ± 0.06*</td>
<td>0.72 ± 0.04</td>
</tr>
<tr>
<td>Concordance between WSI and glass slides</td>
<td>86%</td>
<td></td>
</tr>
<tr>
<td>Concordance and minor discordance between WSI and glass slides</td>
<td>98%</td>
<td></td>
</tr>
</tbody>
</table>

*P=.005 compared to glass slides
Baseline intra/inter-observer variability exists even with glass slides.

Aim is to evaluate the technology, not agreement between pathologists.
  - e.g. Prostate ASAP may have varying pathologist opinions

Therefore, we recommend:
  - **Measure** intra-pathologist diagnostic reproducibility
    i.e. is the pathologist able to reach the same diagnosis with both modalities?
  - **Don’t measure** interobserver variability
    i.e. their diagnosis compared to other pathologists/ experts/ consensus

Comment feedback: 86% agreement
Guideline Statement #9

- Digital and glass slides can be evaluated in random or nonrandom order (as to which is examined first and second) during the validation process.

Grade: Recommendation, Level A
• Some believe that digital slides should be viewed before glass slides - considered the gold standard for making diagnoses.

• However, the order of viewing virtual vs. glass slides has been shown not to affect interpretation (Koch LH et al. Human Pathol 2009; 40:662-7).

• The evidence indicates it can go either way.

Comment feedback: 81% agreement for original random review – Panel made revisions on evidence
Different Outcomes of WSI & Glass Slides with Random or Nonrandom Allocation of Cases

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Allocation of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Random</td>
</tr>
<tr>
<td></td>
<td>WSI</td>
</tr>
<tr>
<td>Accuracy of WSI or glass slides</td>
<td>72%</td>
</tr>
<tr>
<td>Concordance between WSI and glass slides</td>
<td>81%</td>
</tr>
<tr>
<td>Discordance between WSI and glass slides</td>
<td>19%</td>
</tr>
<tr>
<td>Concordance and minor discordance between WSI and glass slides</td>
<td>93%</td>
</tr>
</tbody>
</table>

* P< .001 versus glass slide [nonrandom]
Guideline Statement #10

• A washout period of at least 2 weeks should occur between viewing digital and glass slides.

Grade: Recommendation, Level B
• **Washout period** = time interval between viewing the same case/slide using a different (glass or digital) modality.

• **Important to take into consideration:**
  - Pathologists may recall pathologic images for lengthy periods after reviewing a case.
  - With long washout periods a pathologist’s experience and/or diagnostic criteria could change over time.

**Comment Feedback:** 68% agreement- Panel made revisions according to evidence
### Different Outcomes of WSI and Glass Slides with Different Duration of Washout Periods

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Washout periods for WSI and Glass Slides</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1 wk</td>
</tr>
<tr>
<td>WSI</td>
<td>Glass</td>
</tr>
<tr>
<td>Accuracy of WSI or glass slides</td>
<td>70%</td>
</tr>
<tr>
<td>Concordance between WSI and glass slides</td>
<td>NR</td>
</tr>
<tr>
<td>Discordance between WSI and glass slides</td>
<td>NR</td>
</tr>
<tr>
<td>Concordance and minor discordance between WSI and glass slides</td>
<td>NR</td>
</tr>
</tbody>
</table>
Guideline Statement #11

• The validation process should confirm that all of the material present on a glass slide to be scanned is included in the digital image.

Grade: Expert Opinion
• Accurate digital reproduction of scanned glass slides is required if they are to be used for diagnostic use.

Public comment: 94% agreement
Guideline Statement #12

- Documentation should be maintained recording the method, measurements and final approval of validation for the WSI system to be used in the clinical laboratory.

Grade: Expert Opinion

Comment feedback: 97% agreement
Conclusion

• Validation of WSI is necessary to ensure that a pathologist using this technique to view digitized glass slides can consistently make the same clinical interpretation as they would from viewing the glass slides using a traditional bright field microscope.

• Validation should address both technical and interpretative components, and must be specific for the intended clinical use.

• Ongoing future updates on this topic are planned.