Technical Performance Assessment of Digital Pathology Whole Slide Imaging Devices

Guidance for Industry and Food and Drug Administration Staff

Document issued on: April 20, 2016

The draft of this document was issued on February 25, 2015

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U.S. Department of Health and Human Services
Food and Drug Administration
Center for Devices and Radiological Health

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Preface

Public Comment

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Table of Contents

I.	Introduction	1
II.	Background	2
III.	Scope	2
IV.	Policy	3
IV	(A). Description and Test Methods for Each Component	3
	IV(A)(1). Slide Feeder	5
	IV(A)(1)(a). Description	5
	IV(A)(2). Light Source	5
	IV(A)(2)(a). Description	5
	IV(A)(2)(b). Test Method	6
	IV(A)(3). Imaging Optics	6
	IV(A)(3)(a). Description	6
	IV(A)(3)(b). Test Methods	7
	IV(A)(4). Mechanical Scanner Movement	7
	IV(A)(4)(a). Description	7
	IV(A)(4)(b). Test Method	8
	IV(A)(5). Digital Imaging Sensor	8
	IV(A)(5)(a). Description	8
	IV(A)(5)(b). Test Methods	8
	IV(A)(6). Image Processing Software	9
	IV(A)(6)(a). Description	9
	IV(A)(6)(b). Resources	9
	IV(A)(7). Image Composition	9
	IV(A)(7)(a). Description	9
	IV(A)(7)(b). Test Methods	. 10
	IV(A)(8). Image Files Formats	. 10
	IV(A)(8)(a). Description	. 10
	IV(A)(9). Image Review Manipulation Software	. 11
	IV(A)(9)(a). Description	. 11
	IV(A)(9)(b). Resources	. 11

IV(A)(10). Computer Environment	11
IV(A)(10)(a). Description	11
IV(A)(11). Display	12
IV(A)(11)(a). Description	12
IV(A)(11)(b). Test Methods	12
IV(A)(11)(c). Resources	13
IV(B). System-level Assessment	14
IV(B)(1). Color Reproducibility	15
IV(B)(1)(a). Description	15
IV(B)(1)(b). Test Methods	15
IV(B)(1)(c). Resources	16
IV(B)(2). Spatial Resolution	16
IV(B)(2)(a). Description	16
IV(B)(2)(b). Test Methods	16
IV(B)(3). Focusing Test	16
IV(B)(4). Whole Slide Tissue Coverage	17
IV(B)(4)(a). Description	17
IV(B)(4)(b). Test Method	17
IV(B)(5). Stitching Error	18
IV(B)(5)(a). Description	18
IV(B)(5)(b). Test Methods	18
IV(B)(6). Turnaround Time	19
IV(B)(6)(a). Description	19
IV(C). User Interface	19
IV(C)(1). Description	19
IV(C)(2). Test Methods	19
IV(C)(3). Resources	22
IV(D). Labeling	22
IV(D)(1). Test Methods	23
IV(D)(2). Resources	23
IV(E) Quality Control	23

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Guidance for Industry and Food and Drug Administration Staff

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I. Introduction

FDA is issuing this guidance to provide industry and agency staff with recommendations regarding the technical performance assessment data that should be provided for regulatory evaluation of a digital whole slide imaging (WSI) system. This document does not cover the clinical submission data that may be necessary to support approval or clearance. This document provides our suggestions on how to best characterize the technical aspects that are relevant to WSI performance for their intended use and determine any possible limitations that might affect their safety and effectiveness.

Recent technological advances in digital microscopy, in particular the development of whole slide scanning systems, have accelerated the adoption of digital imaging in pathology, similar to the digital transformation that radiology departments have experienced over the last decade. FDA regulates WSI system manufacturers to help ensure that the images intended for clinical uses are reasonably safe and effective for such purposes. Essential to the regulation of these systems is the understanding of the technical performance of the WSI system and the components in the imaging chain, from image acquisition to image display and their effect on pathologist's diagnostic performance and workflow. Prior to performing non-technical analytical studies (i.e., those using clinical samples) and clinical studies to evaluate a digital imaging system's performance, the manufacturer should first determine the technical characteristics that are relevant to such performance for its intended use and determine any possible limitations

that might affect its safety and effectiveness. This guidance provides recommendations for the assessment of technical characteristics of a WSI device.

FDA's guidance documents, including this guidance, do not establish legally enforceable responsibilities. Instead, guidances describe the Agency's current thinking on a topic and should be viewed only as recommendations, unless specific regulatory or statutory requirements are cited. The use of the word *should* in Agency guidance means that something is suggested or recommended, but not required.

II. Background

For over a hundred years, the reference method for the diagnosis of cancer and many other critical clinical conditions has been histopathological examination of tissues using conventional light microscopy. This process is known as surgical pathology in the United States.

In surgical pathology, patient tissue from surgery, biopsy or autopsy goes through a process that includes dissection, fixation, embedding, and cutting of tissue into very thin slices which are then stained, for example by the hematoxylin and eosin (H&E) protocol, and permanently mounted onto glass slides. The slides are examined by a pathologist under a light microscope by dynamically adjusting the focus and using different magnifications. By integrating their interpretations obtained by microscopic examination of the tissue from all slides pertaining to a case, pathologists arrive at a diagnosis of the case.

WSI refers to the digitization of the stained entire tissue specimen on a glass slide. The glass slide is still prepared and stained just as for conventional light microscopy. Depending on the system used, various magnifications, scanning methodologies, hardware, and software are employed to convert the optical image of the slide into a digital whole slide image. With WSI, the pathologist views the image on a computer monitor rather than through the microscope oculars.

III. Scope

This document provides guidance regarding only the technical performance assessment of WSI systems for regulatory evaluation. WSI systems are defined here as those consisting of (a) an image acquisition subsystem that converts the content of a glass slide into a digital image file, and (b) a workstation environment for viewing the digital images. If not otherwise specified, the term "image" in the context of whole slide imaging refers to a pyramid structure consisting of multiple images at different resolutions. The baseline image has the highest resolution. This guidance is applicable for surgical pathology tasks performed in the anatomic pathology laboratory. It is intended to provide recommendations to industry and FDA staff regarding only the technical performance assessment data needed for the regulatory evaluation of a WSI device. This document is not meant to provide guidance for special stain techniques or

fluorescence imaging or for the non-technical analytical studies (utilizing clinical samples) or pivotal clinical studies necessary to support safety and effectiveness, nor does this guidance alone suffice to demonstrate safety and effectiveness of WSI systems. Interpretation of WSI images on mobile platforms is beyond the scope of this guidance.

IV. Policy

The following subsections of this section describe the technical performance assessment data FDA believes will facilitate the regulatory evaluation of a WSI device.

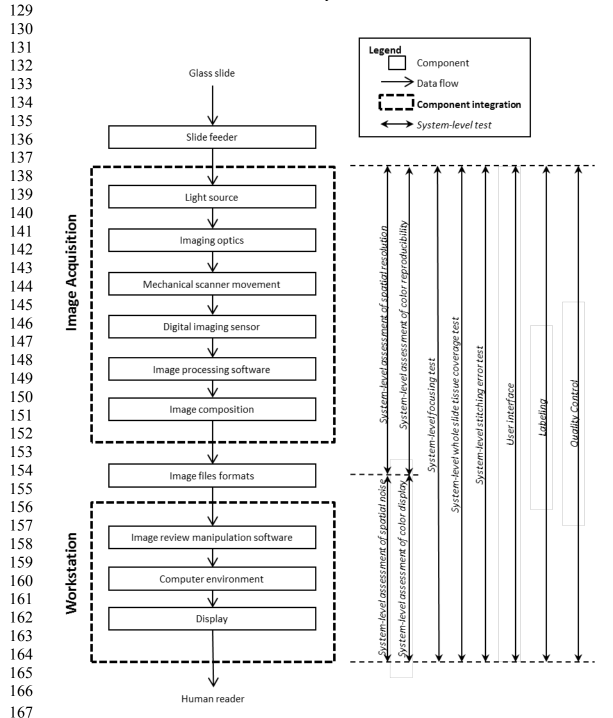
IV(A). Description and Test Methods for Each Component

This subsection details the descriptions and the test methods at the component level that should be included in the technical performance assessment of a WSI device. For purposes of this guidance only, a component is a piece of hardware, software, or a combination of hardware and software that processes the image signals flowing through the imaging chain. The concept of a component is based on the transformation of the image signals. For example, the digital imaging sensor is a hardware device that converts optical signals into digital signals. The image composition component is a software program that stitches sub-images together to form a whole slide image. A component and a physical device need not be in close physical proximity. For example, the light source component and the image optics component are usually tightly coupled within the same device, while the display calibration data is often distributed in both the color profile in the computer environment component and the on-screen display settings in the display component.

The components in a WSI device can be grouped in two subsystems: image acquisition and image display. The image acquisition subsystem digitizes the tissue slide as a digital image file. The image display subsystem converts the digital image file into optical signals for the human reader. In the paradigm of telemedicine, the digital image file can be electronically sent to a remote site for reading, so the image acquisition subsystem and the image display subsystem do not need to be physically coupled. Methods for independently testing the image acquisition and display subsystems are described in Section IV(B).

Sponsors should provide a block diagram of the components found in the WSI system in the premarket submission. A chart indicating the relationship among the components and the test methods utilized for the specific system characterization should also be provided. Diagram 1 on the following page is offered as an example block diagram of typical components found in current WSI systems. The components of a particular WSI system might not include all of those listed in the diagram or may include additional components. Sponsors are encouraged to provide additional diagrams, illustrations, and photographs of their devices as part of their submissions.

Diagram 1: Example block diagram of typical components found in current WSI systems



168	IV(A)(1). Slide Feeder
169	
170	IV(A)(1)(a). Description
171 172	The slide feeder is the mechanism(s) used to introduce the slide(s) to the scanner. For the
173	slide feeder, sponsors should provide the following information, if applicable:
174	Configuration of the slide feed mechanism (a physical description of the
175	equipment)
176 177	 Slide configuration (physical description of the slide (i.e., custom or commercial off the shalf)
178	commercial off-the-shelf)) Number of slides in queue (carrier)
179	
180	 Class of automation (e.g., robotics, pneumatics, etc.) User interaction
181	
182	 Hardware (e.g., loading of slides into carrier) Software (e.g., does the system recognize the number of slides or is this
183	specified by the user)
184	 Feedback (e.g., alarms, notifications, etc.)
185	 Failure Mode and Effects Analysis (FMEA) (including severity,
186	likelihood, mitigations, etc.)
187	
188	IV(A)(2). Light Source
189	1 (11)(2) Digno Source
190	IV(A)(2)(a). Description
191	1 v (r)(2)(u). Description
192	The light source, including the light guide, generates and delivers light to the slide being
193	imaged. The two major components are the lamp and condenser. For the light source,
194	sponsors should provide the following information and specifications, if applicable:
195	• Lamp
196	o Bulb type (e.g., halogen, xenon arc, LED)
197	Manufacturer and model
198	o Wattage
199	 Spectral power distribution
200	 Expected lifetime
201	 Output adjustment control (electrical/electronic/mechanical)
202	Optical filter(s)
203	Type (e.g., heat blocking, polarization, neutral density, diffusing)
204	 Manufacturer and model
205	 Expected intensity variation (coefficient of variation)
206	 Over the duration of scanning a single slide
207	• Over the course of a single workday
208	• Over the lifetime of the device
209	• Expected spectral variation
210	• Over the duration of scanning a single slide
211	• Over the course of a single workday
212	• Over the lifetime of the device
213	 Capability of tracking intensity and spectral degradation with lifetime

214	 Condenser
215	 Illumination format (e.g., Kohler, critical)
216	 Manufacturer and model
217	 Numerical aperture
218	 Focal length
219	 Working distance
220	
221	IV(A)(2)(b). Test Method
222	
223	The following steps should be used to measure the spectral distribution of light incident
224	on the slide. Position the input of a calibrated spectrometer or monochromator at the
225	plane where the slide would be placed, centered on the illumination spot from the
226	condenser. If desired, the light can be coupled into the spectrometer via light guide (e.g.
227	fiber optic cable) or an integrating sphere. The measurement aperture should be at least
228	as large as the anticipated field of view on the slide at the lowest magnification of the
229	imaging optics. The wavelength accuracy and relative spectral efficiency of the
230	spectrometer or monochromator in the wavelength range of 360-830 nm should be
231	calibrated prior to measurements and reported. Plots of the measured spectrum with at
232 233	least 10 nm spectral resolution should be provided, using radiometric units (e.g., spectral irradiance in W/cm ² /nm, spectral radiance in W/sr/cm ² /nm).
234	irradiance in w/cm /mm, spectral radiance in w/si/cm /mm).
235	IV(A)(3). Imaging Optics
	TV (A)(3). Imaging Optics
236	IV(A)(2)(a) Denomination
237238	IV(A)(3)(a). Description
239	The imaging optics comprises the microscope objective and auxiliary lens(es) (e.g., tube
240	lens), which optically transmit an image of the tissue from the slide to the digital image
241	sensor. Sponsors should provide the following information and specifications, if
242	applicable:
243	 Optical schematic with all optical elements identified from slide (object plane) to
244	digital image sensor (image plane)
245	Microscope objective
246	Manufacturer
247	o Type
248	o Magnification
249	o Numerical aperture (NA)
250	o Focal length
251	 Working distance
252	• Auxiliary lens(es)
253	 Manufacturer
254	 Lens type
255	o Focal length
256	• Magnification of imaging optics: ISO 8039:2014 Optics and optical instruments
257	— Microscopes — Magnification
258	
259	

260 261	IV(A)(3)(b).	Test Methods
262 263	Sponsors should conduct the Standards, if applicable:	following tests in conformance with the International
264	· • • •	f imaging optics at image plane per ISO 13653:1996 Optics
265	and optical instrumen	nts – General optical test methods - Measurement of relative
266	irradiance in the imag	2 4
267		39:2008 Optics and photonics — Quality evaluation of
268	÷ •	termination of distortion
269		s per ISO 15795:2002 Optics and optical instruments —
270		Coptical systems — Assessing the image quality degradation
271	due to chromatic abe	rrations
272		
273	IV(A)(4). Mech	nanical Scanner Movement
274	TY/AN/AN/	D 14
275	IV(A)(4)(a).	Description
276277	The machanical gapmar add	ranges the physical characteristics of the stage upon which
278		resses the physical characteristics of the stage upon which
279		e key components include stage configuration, movement, n is relevant whether it is only the stage that is moving and
280		f there is movement on all axes. For the mechanical scanner
281	1	following information and specifications, if applicable:
282	-	stage (a physical description of the stage)
283	o Stage size	stage (a physical description of the stage)
284		cturer and model number
285		l (e.g., anodized aluminum)
286		ixis or multiple stacked linear stages (manufacturer and
287	model number	
288		s or ways (e.g., bearings)
289		ion mechanism (slide holder)
290	-	t of the stage (e.g., stepper motor, servomotor, piezomotor,
291		t, ball-screw, lead-screw, etc.)
292	, 1	solution for XY-axes
293	 Movement in 	
294	 Speed range 	
295	 Travel distance 	ee
296	 Maximum sca 	nning area
297	 Localization a 	and reading of bar code labels
298	 Control of movement 	of the stage
299	 Open or close 	d loop operation
300	 Positional acc 	uracy (calibration) and repeatability
301	Lost m	notion compensation (e.g., backlash)
302		rol (e.g., joystick) for single-slide, non-batch mode
303	 Selection of an 	rea to be scanned (in accordance to image composition
304	software)	
305	whole	slide

306	 automatically determined area with tissue content
307	• Failure Mode and Effects Analysis (FMEA) (including severity, likelihood,
308	mitigations, etc.)
309	
310	IV(A)(4)(b). Test Method
311 312	Sponsors should demonstrate the mechanical performance of the stage with respect to
313	positional repeatability and accuracy on all relevant axes, in accordance with ISO 230-
314	2:2014 Test code for machine tools—Part 2: Determination of accuracy and
315	repeatability of positioning numerically controlled axes.
316	
317	IV(A)(5). Digital Imaging Sensor
318	
319	IV(A)(5)(a). Description
320	
321	The digital image sensor is an array of photosensitive elements (pixels) that convert the
322	optical signals of the slide to digital signals, which consist of a set of values
323	corresponding to the brightness and color at each point in the optical image. Please
324	provide the following information and specifications:
325	 Sensor type (e.g., CMOS, CCD) and manufacturer
326	 Pixel information/specifications
327	 Number and dimensions of pixels
328	 Design of color filter array
329	 Configuration of color filter array
330	 Spectral transmittance of color filter mask
331	• Responsivity specifications
332	 Relative response versus wavelength
333	Continuo de la continuación d
334	 Spatial uniformity
335	 Noise specifications
336	 Dark current level (electrons per second)
337	 Read noise (electrons)
338	 Readout rate (e.g., pixels per second, frames per second)
339	• Digital output format (e.g., bits per pixel, bits per color channel)
340	
341	IV(A)(5)(b). Test Methods
342	
343	Sponsors should conduct the following tests in conformance with the corresponding
344	International Standards, if applicable:
345	
346	• Opto-electronic conversion function per ISO 14524:2009 <i>Photography</i> —
347	Electronic still-picture cameras — Methods for measuring optoelectronic
348	conversion functions (OECFs)
349	Noise measurements per ISO 15739:2013 Photography — Electronic still-picture
350	imaging — Noise measurements
351	

352	IV(A)(6). Image Processing Software
353	
354	IV(A)(6)(a). Description
355	
356	Image processing software refers to the embedded software components of the image
357	acquisition device. It typically includes control algorithms for image capture and
358	processing algorithms for raw data conversion into the digital image file. Sponsors
359	should provide the following information and specifications, if applicable:
360	• Exposure control
361	• White balance
362	• Color correction
363	• Sub-sampling
364	Pixel-offset correction
365	Pixel-gain or flat-field correction
366	Pixel-defect correction
367	W(A)(C)(b) Description
368 369	IV(A)(6)(b). Resources
370	See the guidance entitled "Guidance for the Content of Premarket Submissions for
371	Software Contained in Medical Devices"
372	(http://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocument
373	s/ucm089543.htm) for the information that should be provided.
374	
375	IV(A)(7). Image Composition
376	
377	IV(A)(7)(a). Description
378	
379	Image composition is a step present in systems that produce whole slide images as
380	opposed to individual fields of view. Whole slide scanning is typically performed in
381	accordance with the positioning of a stage that moves in submicron steps. At each
382	location of the stage movement, an image of the field of view is acquired. Images can be
383	acquired with a degree of overlapping (redundancy) between them to avoid gaps in data
384	collection. Images can also be acquired at different depths of focus followed by the
385	application of focusing algorithms. At the end of this process, all acquired images are
386	combined (stitched) together to create a composite high resolution image. There are a
387 388	number of features that can affect this process, and they are listed below. Sponsors
389	should provide a description of these features, if applicable: • Scanning method
390	 Scanning method Single objective or multiple miniature objectives in an array pattern
390	 Single objective of multiple inimature objectives in an array pattern Scanning pattern: square matrix acquisition (tiling), line scanning, etc.
392	 Overlap between scanned regions
393	 Merging algorithms that stitch the aligned images together into a
394	composite image file. Such algorithms may employ functions to align
395	adjacent fields of view in accordance to the scanning pattern, overlap, etc.

396 397 398 399 400 401 402 403	 Automatic background correction functions to eliminate the effect of non-uniformities in the microscope's illumination and image merging procedure. These non-uniformities if not corrected might create visible borders (seams and stitch lines) between the adjacent fields of view. Scanning speed: time to scan the whole slide. This time is dependent on selected magnification, and the amount of tissue on the glass slide. Number of planes at the Z-axis to be digitized (stack depth)
404	IV(A)(7)(b). Test Methods
405	
406	Testing for image composition can be performed on a system level using special
407	calibration slides (such as grid patterns) that can test for line uniformity and focus
408	quality. Sponsors should provide the following outputs for these tests, if applicable:
409	 Images of digitized calibration slides
410	 Analysis of focus quality metrics
411	 Analysis of coverage of the image acquisition for the entire tissue slide
412	
413	IV(A)(8). Image Files Formats
414	
415	IV(A)(8)(a). Description
416	
417	The final result from image acquisition can be a whole slide image consisting of a stack
418	of all acquired fields of view and magnifications during WSI. The complete digitized
419	image file usually occupies between 1-20 gigabytes of storage space depending on the
420 421	sample and the magnification of the objective lens used. Images can then be stored in a
421	number of ways and formats. Sponsors should provide the following information:
423	• Compression method (e.g., the wavelet-based JPEG2000 compression standard or TIFF)
424	 Compression ratio: ratio of uncompressed to compressed file size. This metric
425	should be provided along with descriptive information on the data it was
426	measured from, since compression ratio is dependent on the content of the data
427	applied to.
428	 Compression type: lossless or lossy compression
429	• File format: can be formats easily accessible with public domain software such as
430	JPEG or TIFF, or can be proprietary formats only accessible with specific vendor
431	viewers. The file format depends on the file organization and related use.
432	• For systems that interact with DICOM-compliant software and hardware,
433	sponsors should provide a DICOM compatibility report.
434	• File organization:
435	 Single file with multi-resolution information (pyramidal organization)
436	 Stack of files at different magnifications
437	
438	
439	

440	IV(A)(9). Image Review Manipulation Software
441	
442	IV(A)(9)(a). Description
443	
444	For the image review manipulation software, sponsors should provide the following
445	information, describing software features, if applicable.
446	 Continuous panning (moving in x-y space) and pre-fetching (buffering adjacent
447	images to speed up panning time)
448	 Continuous zooming (magnification)
449	 Discrete Z-axis displacement
450	 Ability to compare multiple slides simultaneously on multiple windows
451	 Ability to perform annotations
452	Image enhancement such as sharpening functions
453	• Color manipulation, including color profile, white balance, color histogram
454	manipulation, and color filters
455	 Annotation tools
456	 Tracking of visited areas and annotations
457	 Digital bookmarks (revisit selected regions of interest)
458	 Virtual "multihead microscope" (this is when multiple pathologists
459	simultaneously review the same areas remotely)
460	
461	IV(A)(9)(b). Resources
462	
463	See the guidance entitled "Guidance for the Content of Premarket Submissions for
464	Software Contained in Medical Devices"
465	(http://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocument
466	s/ucm089543.htm) for additional information on this subject.
467	
468	IV(A)(10). Computer Environment
469	
470	IV(A)(10)(a). Description
471	
472	Computer environment refers to the workstation, including both hardware and software
473	components, that retrieves the digital image file and drives the display for the user to
474 475	review the images. Sponsors should provide the following information and
475	specifications, if applicable:
476	Computer hardware Or grating gystem
477	Operating system Crapbias and
478	• Graphics card
479	Graphics card driver
480	Color management settings
481	• Color profile
482 483	 Display interface (e.g., DVI or DisplayPort)
41 X 🔞	

484	IV(A)(11). Display
485	
486	IV(A)(11)(a). Description
487	
488	The final stage of a WSI system is the display component that presents the scanned image
489	to the pathologists for reading. Technically, display refers to the optoelectronic device
490	that converts the digital image signals in the RGB space into optical image signals. For
491	the display, sponsors should provide the following information and specifications, if
492	applicable:
493	• Technological characteristics of the display device (e.g., in-plane switching LCD
494	panel with TFT active-matrix array with fluorescent backlight)
495	Physical size of the viewable area and aspect ratio
496	• For transmissive displays, backlight type and properties including temporal,
497	spatial, and spectral characteristics
498	• Frame rate and refresh rate
499	• Pixel array, pitch, pixel aperture ratio and subpixel matrix scheme (e.g., chevron,
500	RGBW)
501	• Subpixel driving to improve grayscale resolution (e.g., spatial and temporal
502	dithering)
503	• Supported color spaces
504	Display Interface
505	• User controls of brightness, contrast, gamma, color space, power-saving options,
506	etc. via the on-screen display (OSD) menu
507	• Ambient light adaptation including the ambient light sensing method,
508	instrumentation, and software tool description
509	• Touch screen technology including method, functionality, and any calibration or
510 511	periodical re-tuning requirements
512	 Color calibration tools (sensor hardware and associated software), color profile, and method for color management
513	 Frequency and nature of quality-control tests to be performed by the user and/or
514	the physicist with associated action limits.
314	the physicist with associated action mints.
515	
516	IV(A)(11)(b). Test Methods
517	
518	• <i>User controls:</i> Modes and settings of the display undergoing testing should be
519	specified, including brightness, contrast, gamma, white point, color space, etc.
520	See 2.1 Modified-Performance Modes, IDMS 1.03.
521	• Spatial resolution: Measurements of the transfer of information from the image
522	data to the luminance fields at different spatial frequencies of interest typically
523	done by reporting the modulation transfer function. Non-isotropic resolution
524	properties should be characterized properly by providing two-dimensional
525	measurements or measurements along at least two representative axes. See 7.7
526	Effective Resolution, IDMS 1.03.

• *Pixel defects (count and map):* Measurements (counts) and location of pixel defects. This is typically provided as a tolerance limit. Pixel defects can interfere with the visibility of small details in medical images. See 7.6 *Defective Pixels*, *IDMS 1.03*.

- *Artifacts:* Evaluate for image artifacts such as ghosting and/or image sticking from displaying a fixed test pattern for a period of time. See 4.6 *Artifacts and Irregularities*, *IDMS 1.03*.
- *Temporal response:* Measurements of the temporal behavior of the display in responding to changes in image values from frame to frame. Since these transitions are typically not symmetric, rise and fall time constants are needed to characterize the system. See *10.2.3 Gray-to-Gray Response Time, IDMS 1.03*.
- Maximum and minimum luminance (achievable and recommended): Measurements of the maximum and minimum luminance that the device outputs as used in the application under recommended conditions and the achievable values if the device is set to expand the range to the limit. See 2.4 Vantage-Point Suite of Measurement, IDMS 1.03.
- *Grayscale:* Measurements of the mapping between image values and the luminance. See *6.1 Grayscale, IDMS 1.03*.
- Luminance uniformity and Mura test: Measurements of the uniformity of the luminance across the display screen. See 8.1.2 Sampled Vantage-Point Uniformity and 8.2.3 Mura Analysis, IDMS 1.03.
- Stability of luminance and chromaticity response with temperature and lifetime
- *Bidirectional reflection distribution function:* Measurements of the reflection coefficients of the display device. Specular and diffuse reflection coefficients can be used as surrogates for the full bidirectional reflection distribution function. See 11.12 Diagnostic: Characterizing Hemisphere Uniformity, IDMS 1.03.
- *Gray Tracking:* Chromaticity at different luminance levels as indicated by the color coordinates in an appropriate units system (e.g., CIE *u'v'*). See *AAPM Task Group 196 Report*.
- *Color scale:* Color coordinates of primary and secondary colors as a function of the digital driving level and their additivity. See 6. *Gray- and Color-Scale Measurement and 5.4 Color-Signal White, IDMS 1.03*.
- Color gamut volume: See 5.31 Volume-Color-Reproduction Capability, IDMS 1.03.

IV(A)(11)(c). Resources

Those interested in learning more about these types of display considerations should consider reading:

- IDMS 1.03 Information Display Measurements Standard Version 1.03, International Committee for Display Metrology, Society for Information Display, www.icdm-sid.org
- E. Samei, A. Badano, D. Chakraborty, K. Compton, C. Cornelius, K. Corrigan, M. J. Flynn, B. Hemminger, N. Hangiandreou, J. Johnson, M. Moxley, W.

Pavlicek, H. Roehrig, L. Rutz, J. Shepard, R. Uzenoff, J. Wang, and C. Willis,
Assessment of display performance for medical imaging systems, Report of the
American Association of Physicists in Medicine (AAPM) Task Group 18,
Technical Report, AAPM (April 2005).

• IEC 62563-1:2009, Medical electrical equipment – Medical image display systems – Part 1: Evaluation methods

• Amendment 1 to IEC 62563-1: *Medical image display systems – Part 1: Evaluation methods*

- The guidance entitled "Guidance for Industry and FDA Staff: Display Accessories for Full-Field Digital Mammography Systems-Premarket Notification (510(k)) Submissions"
 - (http://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/ucm107549.htm).

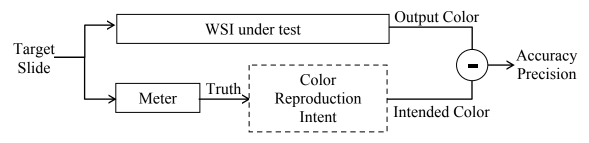
IV(B). System-level Assessment

This subsection details the test methods at the system level that should be included in the technical performance assessment of a WSI device. In this guidance, *system* refers to a series of consecutive components in the imaging chain with clearly defined, measureable input and output. For example, a system-level test can be designed for the image acquisition subsystem, the image display subsystem, or a combination of both. The goal of system-level tests is to assess the composite performance of a series of consecutive components in the imaging chain. System-level tests should be conducted when the component-level tests are either unfeasible or unable to capture the interplay between components.

The common framework of the system-level tests described in this section is to compare the system under test with an ideal system based on the same input, and then report the difference between their outputs quantitatively. Designing such a system-level test typically involves the following steps: (1) define the scope of the system and its input and output, (2) define the input, which in most cases is a test target or phantom, (3) measure the input to establish the ground truth that would be generated by an ideal system, (4) measure the output of the system under test, and (5) calculate the errors between the truth and the output with a quantitative metric. The framework of a typical system-level test is shown in Diagram 2. Notice that the *ideal system* is a hypothetical device that generates the perfect output with respect to the objective of the test such as color or focus. The purpose of the ideal system is to define the intended behavior of the system under test. The ideal system does not need to be implemented. Instead, the ideal system should be simulated by a test method that establishes the truth of the input phantom.

619	Diagram 2: Framework of a typical system-level test.
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621	System under test —>Output —
622	$ \begin{array}{c} \text{Input} \\ \text{(Phantom)} \end{array} $ Error
623	Ideal system Truth
624	Ideal System
625	
626	IV(B)(1). Color Reproducibility
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628	IV(B)(1)(a). Description
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630	Color reproducibility is one of the key characteristics of a WSI system. The color
631	characteristics are determined by every component in the imaging chain. Therefore, the
632	color characteristics might be best evaluated at the system level. Color reproducibility
633	indicates the accuracy and precision of the color transformation from the tissue sample on
634	the slide to the image on the display. The colors of the tissue specimen should be
635	accurately and precisely reproduced on the display based on the color reproduction intent,
636	which should be clearly defined and justified by the sponsor.
637	
638	IV(B)(1)(b). Test Methods
639	The WCI and an about the first decided and a side of the standard and all the standard and a side of the side of the standard and a side of the standard and a side of the side o
640 641	The WSI system should be tested with a target slide. The target slide should contain a set of measurable and representative color patches. Ideally the color patches should have
642	similar spectral characteristics to stained tissue. The color patches should include a
643	grayscale ramp for evaluating the grayscale response. The truth of the color patches
644	should be measured with proper apparatuses separately.
645	should be measured with proper apparatuses separately.
646	For each color patch, the intended color (i.e., the expected output color based on the color
647	reproduction intent defined by the Sponsor) should be calculated based on the truth of the
648	color patches.
649	r
650	The target slide should be scanned and displayed by the WSI system. The output color of
651	each color patch should be measured from the display.
652	
653	The three datasets – truth, intended color, and output color – should be compared and
654	analyzed. The sponsor should provide a rationale if the intended color is different from
655	the truth.
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Diagram 3: Framework of the system-level color reproducibility test.



IV(B)(1)(c). Resources

Useful references on the subject of color reproducibility can be found at the International Color Consortium website http://www.color.org.

IV(B)(2). Spatial Resolution

IV(B)(2)(a). Description

Spatial resolution is another key characteristic of a WSI system. The goal of this system-level test is to evaluate the composite optical performance of all components in the image acquisition phase (i.e., from slide to digital image file).

IV(B)(2)(b). Test Methods

The following test is recommended for assessing spatial resolution of the image acquisition phase:

Resolution and spatial frequency response: ISO 12233:2014(E) — Photography
 Electronic still picture imaging — Resolution and spatial frequency responses.

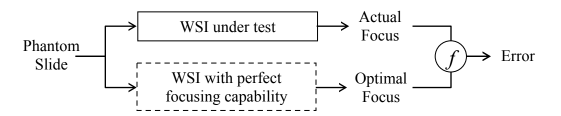
IV(B)(3). Focusing Test

- The quality of focus in WSI can be affected by a number of inter-related factors, including the scanning method and approaches for constructing a focus map. Due to a trade-off between the number of focus points and the overall speed of the scanning process, focusing is typically based on a sample of focus points, determined automatically (auto-focus) or manually by the user. Since tissue can have uneven depth, auto-focus algorithms are needed to detect and adjust for different depths of focus.
- Data demonstrating that the focus quality is acceptable, even in the presence of uneven tissue, should be provided. Such data with proper justification could be derived from a phantom study, from clinical data, or both in a complementary fashion. The technology of phantom construction for testing focus is under development and this guidance will be updated as such technologies become available. Sponsors could attempt to build their own phantoms for testing depth

of focus for their device. Alternatively, sponsors could provide experimental data using clinical tissue slides. Sampling of cases for such an experiment should be enriched for uneven tissue cases within a range representative of typical laboratory output. Alternative approaches for assessing the focus quality of a WSI will be considered along with proper justification. In addition, the following specifications should be provided, if applicable:

- Focus method: auto-focus for high-throughput or user-operated focus points
- Instructions for the selection of manual focus points (if applicable), including number of focus points and location in relation to a tissue sample
- Metrics used to evaluate focusing and description of methods to extract them
- Methods for constructing focus map from sample focus points

Diagram 4: Framework of the system-level focusing test.



IV(B)(4). Whole Slide Tissue Coverage

IV(B)(4)(a). Description

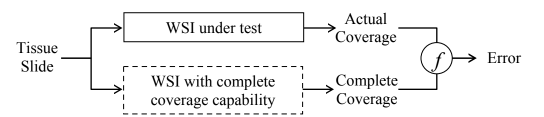
During the scan phase, WSI systems usually skip blank areas where tissue is absent in order to reduce scan time and file size. The purpose of the whole slide tissue coverage test is to demonstrate that all of the tissue specimen on the glass slide is included in the digital image file.

IV(B)(4)(b). Test Method

Sponsors should include a test that demonstrates the completeness of the tissue coverage. Sponsors should describe the test method and include the following items:

- Selection of the input tissue slide
- How to determine the complete coverage of the input tissue slide
- How to measure the actual coverage of the WSI output
- Calculate the ratio of the actual to complete coverage

 Diagram 5: Framework of the system-level whole slide tissue coverage test



Stitching Error IV(B)(5).

Description IV(B)(5)(a).

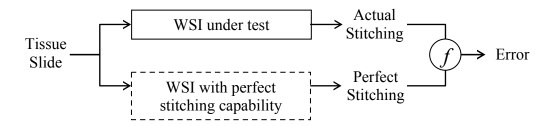
Stitching is the technique that enables a WSI system to combine thousands of sub-images into a single whole-slide image. Although during the scanning process a certain amount of overlapping between adjacent sub-images is maintained for alignment purposes, successful stitching relies on the texture present in the overlapped area. When the stitching algorithm fails to align two sub-images seamlessly, the error may or may not be perceivable by the human reader depending on whether noticeable stitching artifacts are generated. Therefore, a system-level test should be conducted when assessing the stitching quality of the WSI system.

IV(B)(5)(b). **Test Methods**

Sponsors should include a test that evaluates the stitching errors and include the following items:

- Selection of the input test slide
- Method for sampling of the stitching boundaries where stitching errors might occur
- How to determine the ideal stitching as the ground truth
 - o For example, the region of the stitching boundaries can be re-imaged in one shot such that there is no stitching artifact.
- How to evaluate quality of the actual stitching based on the perfect stitching
 - o For example, compare the image of stitching boundaries with the perfect one that does not have stitching artifact. The difference between these two images can be used as a figure of merit of the stitching quality.

Diagram 6: Framework of the system-level stitching error test



IV(B)(6). Turnaround Time

IV(B)(6)(a). Description

Turnaround time is the time required by the WSI system to execute a particular user operation such as panning/zooming where the software and I/O (input/output) devices retrieve image data, execute the computation, and refresh the image on the display. The turnaround time starts when the user enters a command via a keyboard stroke or a mouse click/movement and finishes when the image is completely updated on the display. Turnaround time is important for a WSI system when fast and repetitive panning operations are performed during a search task, which is delay-free in an optical microscope. Prolonged, unpredictable turnaround time may impact the user's diagnostic performance. The user interface should properly prompt the user when the operation is incomplete and the requested image is not available. The turnaround time may vary greatly depending on the user-requested operation, image content, data size/location, computer workload, display size, etc. The sponsor should report the typical turnaround time as well as the test method and test conditions.

IV(C). User Interface

IV(C)(1). Description

The user interface covers all components and accessories of the WSI system with which users interact while loading the slides and acquiring, manipulating, and reviewing the images. It also includes preparing the system for use (e.g., unpacking, set up, calibration), and performing maintenance. Elements of the user interface have been noted in many of the preceding sections and include two broad categories:

- Options through which the user operates the WSI system, such as:
- 830 Software menu options (e.g., scanning parameters)
 - o Physical controls (e.g., clips on the slide feeder)
 - o Connectors and connections (e.g., cables connecting system components)

• Information presented to the user through

Visual displays (e.g., scanned image, software menus)
 Sounds (e.g., tone played when scanning completed)

Instructions (e.g., software users' manual)

Test Methods

o Labels

IV(C)(2).

It is recommended that the analysis to identify the use-related hazards of the WSI system include the consideration of use errors involving failure to acquire, perceive, read, interpret, and act on information from the WSI system correctly or at all and the harm that could be caused by such errors. A human factors/usability validation test should be performed to demonstrate that representative users of the WSI system can perform essential tasks and those critical to safety under simulated use conditions.

When selecting participants for validation testing, sponsors should carefully consider user capabilities and expectations that could potentially impact the safe and effective use of the WSI system. Examples of items that should be considered, if applicable, include visual acuity and type of vision correction and the impact of expectations formed from prior experience with other systems (e.g., optical microscope).

When selecting the critical tasks to be evaluated, sponsors should incorporate all known use related errors and problems from similar devices (devices having similar technological characteristics and indications for use) into the validation testing. Consideration also should be given to whether task performance changes over time, and if test duration needs to account for user fatigue. Examples might include a user altering a task sequence in response to fatigue from repetitive image selection and manipulation with mouse or keyboard.

When creating the simulated use conditions for validation testing, special consideration should be given to the location of the WSI system primary workstation, its components, their arrangement and how their locations affect user performance. Examples of location considerations might include multiple monitors, a monitor with sub-optimal display settings, or glare on a monitor from indoor lighting.

A human factors/usability validation test report should generally include the information found in Table 1.

Table 1: Items a Human Factors/Usability Validation Test Report Should Include

Section	Contents
1	 Intended device users, uses, use environments, and training Intended user population(s) and critical differences in capabilities between multiple user populations Intended uses and operational contexts of use Use environments and key considerations Training intended for users and provided to test participants
2	 Device user interface Graphical depiction (drawing or photograph) of device user interface Verbal description of device user interface
3	 Summary of known use problems Known problems with previous models Known problems with similar devices

Design modifications implemented in response to user difficulties 4 User task selection, characterization and prioritization Risk analysis methods Use-related hazardous situation and risk summary • Critical tasks identified and included in HFE/UE validation tests Summary of formative evaluations 5 • Evaluation methods • Key results and design modifications implemented • Key findings that informed the HFE/UE validation testing protocol 6 Validation testing Rationale for test type selected (i.e., simulated use or clinical evaluation) Number and type of test participants and rationale for how they represent the intended user populations Test goals, critical tasks and use scenarios studied Technique for capturing unanticipated use errors Definition of performance failures Test results: Number of device uses, success and failure occurrences Subjective assessment by test participants of any critical task failures and difficulties Description and analysis of all task failures, implications for additional risk mitigation 7 Conclusion A statement to the effect that "The <device name/model> has been found to be reasonably safe and effective for the intended users, uses and use environments" should be included under the following conditions: The methods and results described in the preceding sections support this conclusion. Any residual risk that remains after the validation testing would not be further reduced by modifications of design of the user interface (including any accessories and the Instructions for Use (IFU)), is not needed, and is outweighed by the benefits that

may be derived from the device's use.

Recommended methods for performing a human factors/usability validation test are described in the resources listed in section IV(C)(3) entitled "Resources" directly below. The goal of testing is to assure that users can operate the WSI system successfully for the intended uses without negative clinical consequences to the patient and that potential use errors or failures have been eliminated or reduced.

IV(C)(3). Resources

FDA recognizes standards published by national and international organizations that apply human factors engineering/usability engineering (HFE/UE) principles to device design and testing. The recognized standards listed below provide suggestions on conducting an analysis of use-related hazards and a human factors/usability validation test to assess the safety and effectiveness of the final device design.

- ISO 14971:2007, Medical Devices Application of Risk Management to Medical Devices: Provides systematic process to manage the risks associated with the use of medical devices.
- AAMI/ANSI HE75:2009, *Human Factors Engineering Design of Medical Devices*: Comprehensive reference of recommended practices related to human factors design principles for medical devices.
- IEC 62366-1:2015, *Medical devices Application of usability engineering to medical devices:* Describes the process to conduct medical device usability testing and incorporate results into a risk management plan.

In addition, FDA has published guidance with human factors related recommendations to assist manufacturers and facilitate premarket review. The guidance entitled "Guidance for the Content of Premarket Submissions for Software Contained in Medical Devices" (http://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/ucm089543.htm). This guidance document provides recommendations to industry regarding premarket submissions for software devices, including stand-alone software applications and hardware-based devices that incorporate software. It includes test methods to assure that the software conforms to the needs of the user and to check for proper operation of the software in its actual or simulated use environment.

IV(D). Labeling

The premarket application must include labeling in sufficient detail to satisfy the requirements of 21 CFR Part 801 and 21 CFR 809.10. The labeling includes supplementary information necessary to use and care for the WSI system such as instruction books or direction sheets and software user manuals.

Although instructions, labeling, and training can influence users to use devices safely and effectively, they should not be the primary strategy used to control risk. Modification of the user interface design is a more effective approach to mitigate use-related hazards.

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IV(D)(1). **Test Methods**

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It is recommended that studies on labeling and training be conducted separately from other human factors/usability validation testing. Human factors/usability validation testing should be conducted with the final version of the labeling and related materials. Timing and content of training should be consistent with that expected of actual users.

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IV(D)(2). Resources

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FDA has published several guidance documents on labeling to facilitate premarket review and assist manufacturers.

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The guidance entitled "Labeling - Regulatory Requirements for Medical Devices" (http://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/ GuidanceDocuments/UCM095308.pdf).

932 933 934 This publication covers labeling issues that device manufacturers, reconditioners, repackers, and relabelers should consider when a product requires labeling. Labeling includes adequate instructions for use, servicing instructions, adequate warnings against uses that may be dangerous to health, or information that may be necessary for the protection of users.

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• The guidance entitled "Device Labeling Guidance #G91-1 (blue book memo)" (http://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/GuidanceD ocuments/ucm081368.htm).

This guidance is intended to ensure the adequacy of, and consistency in device labeling information. It is intended for use by industry in preparing device labeling.

944 945 946 • The guidance entitled "Human Factors Principles for Medical Device Labeling" (http://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/ GuidanceDocuments/UCM095300.pdf).

947 948 949 This report presents the principles of instruction, human factors, and cognitive psychology that are involved in designing effective labeling for medical devices.

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IV(E). Quality Control

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Sponsors should provide information on the quality control procedures, including frequency and testing methods to be performed by the laboratory technologists and/or field engineers with associated quantitative action limits. Discussions of tests for constancy should include discussions of the slide feeder and scanning mechanisms, coverage of the entire tissue slide, the bar code reader, the light source, the imaging sensor device, and the calibrations at the component and system level. A detailed quality control manual should be provided.