This guidance was written prior to the February 27, 1997 implementation of FDA's Good Guidance Practices, GGP's. It does not create or confer rights for or on any person and does not operate to bind FDA or the public. An alternative approach may be used if such approach satisfies the requirements of the applicable statute, regulations, or both. This guidance will be updated in the next revision to include the standard elements of GGP's.

## POINTS TO CONSIDER FOR CERVICAL CYTOLOGY DEVICES

## Introduction

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This points-to-consider document supplements existing FDA guidance for premarket submission of in vitro diagnostic devices for approval or clearance. The information requested is intended to be comprehensive but may not be all inclusive. The emphasis is on FDA's particular concerns in the review of cytology devices for gynecologic specimens. These products include:

13		-	
14		1.	Collection devices for the initial cervicovaginal specimen [spatulas, brushes,
15			etc.] (regulated under 21CFR 884.4530, class II)
16			
17		2.	Devices for making cell suspensions for thin layer or monolayer slides
18			(unclassified)
19			
20		3.	Staining devices (regulated under 21CFR 864.3800, exempt)
21			
22		4.	Computer-assisted cell-locator devices (regulated under 21CFR 864.5260, class
23			II)
24			
25		5.	Semiautomated and automated computer-assisted image analyzers (unclassified)
26			
27		6.	Any other devices used in preparation, reading and interpretation of
28			gynecologic cytology specimens. (These may be regulated or unclassified.)
29	<b>D</b> .		e in the standard and the standard and the standard and the standard and the standard
30	Desig	ners of	t gynecologic cytology devices have study design and method validation
31	cnaile	enges u	hat are not found in other in vitro diagnostic devices. Some examples are:
.52	*	0	50 william Day tests are performed each user and shout 2 million are discussed
33	Ŧ	Over	50 million Pap tests are performed each year and about 2 million are diagnosed
25		as au	s consequences in the detection rate of the tests
35		majo	consequences in the detection rate of the tests.
30	*	The	loss of even a few cells prior to making a final slide may be critical
29		The .	loss of even a few certs prior to making a mai side may be endeal.
30	*	Ther	e are no animal models or sources of cellular materials to serve as positive or
40		nega	tive internal controls for each nation sample.
41		nogu	the mount control for and haven purched
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1 \* Repeat cervical cytology samples may be unreliable until the cervical epithelium has a 2 sufficient time interval to regenerate, usually at least 4 to 6 weeks. 3 4 \* Cytologic diagnoses are dependent on multiple pre-analytic factors including sample 5 collection and preparation even before the demanding microscopic examination. 6 7 \* Cytologic diagnoses are based on qualitative criteria for image interpretation that may 8 be interpreted differently among individual observers and among laboratories. Some 9 degree of subjectivity is unavoidable. 10 11 FDA encourages sponsors to consult with the Division of Clinical Laboratory Devices 12 (DCLD) for guidance and review of protocols prior to submission of a formal application, preferably prior to commencement of clinical studies. If the studies in the submission 13 14 document the safety and effectiveness of the device, the FDA clearance/approval process can proceed in the most timely manner within the limits of the queue of submissions. 15 16 17 Minimal data required: 18 I. 19 **Device Description** 20 21 Provide the following information: 22 23 Intended use Α. 24 25 A description of the clinical intended use including the clinical disorders for which the device is used, the scientific basis for the disorder(s), the clinical 26 significance of the procedure or test results, the risk-benefit issues for use of 27 the device, and the clinical utility. 28 29 30 Β. Principles of the procedure 31 32 A description of the principles of the procedure or test methodology including 33 what the device does, how it is to be used, and who will use the device. Is the 34 device a component of a system or used as a stand-alone device? Is the device for initial diagnosis of slides, confirmation of the diagnosis, screening, or 35 monitoring? Define who will use this device; cytotechnologists, pathologists, 36 clinicians, gynecologists, laypersons, or a combination of users. 37 38 39 С. Device components 40 A description of the components that are provided with the device. Supply 41 42 instructions for acquiring any components not included with the device. 2

	Points	to Consi	ider: Cervical Cytology Devices, Version 7/25/94
1 2		D.	Manufacturing process.
- 3 4 5 6 7			Documentation that the device is manufactured following FDA Good Manufacturing Practices including product design, quality control, consistency of manufactured product with the original submission product, stability of end product, and any other applicable feature of the device.
8	П.	Prot	ocols
10 11		Α.	General considerations for protocols
12 13			Submit the entire protocol and study plan to FDA for comment.
14 15 16 17 18			Provide a written protocol complete for all phases of the study that is applicable to all study sites before specimen collection/selection begins. It is the responsibility of the sponsor to ensure adherence to the protocol at all study sites.
19 20 21 22 23			Consult with a biostatistician during the initial planning stages of the protocol and during data analysis. <i>The most common reason for premarket submissions</i> <i>to fail is poor planning of the protocol.</i> Once a study is completed, it is almost impossible to make a poorly planned study acceptable.
24 25 26			Ensure that the study has a defined hypothesis and that the study design has the power to test this hypothesis.
27 28 29 30 31 32			Once the study begins, justify and document any and all changes to the protocol. All changes <u>must</u> be written into the protocol. Inform FDA of proposed protocol changes and what effects these changes will have on data interpretation. Discuss these points with the biostatistician prior to implementation.
33 34			Collect data for each intended specimen type or target population.
35 36 37			Collect performance data using a final production model device and not a prototype.
38 39 40 41 42			Appoint a study coordinator to oversee <u>all</u> aspects of the protocol study. This person must have knowledge of all details of the study and serves to protect the integrity of the data.

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1 2 3 4 5 6 7 8		Most study designs will compare a new cytology device to the conventional manually prepared and manual microscopy-read method. In some cases, the new cytology devices may yield test results that were not detected by the conventional PAP test. All abnormal diagnoses on slides that were previously diagnosed as negative must be reported to the laboratory. This includes ASCUS and above, AGCUS and above, etc. The sponsor has the responsibility of monitoring and assuring appropriate patient follow-up. A confirmatory test must be performed according to the Bethesda System (TBS) guidelines.
10	В.	Pre-Clinical Studies
12 13		Define and test for the following performance characteristics:
14 15 16		Accuracy (lack of bias) Comparison to a gold standard
17 18 19 20		Precision (lack of random error) Reproducibility (inter-observer, -laboratory, -instrument, intra-observer,- instrument)
20 21 22		Percent agreement, adjusted
23 24		Morphologic criteria
25 26 27 28 29		Provide data to demonstrate the ability of the device to preserve morphological structure of the cells in comparison to the conventional smear; to capture adequate representation of endocervical components; and to indicate the presence of inflammation and infectious agents.
30 31 32		FDA recommends adherence to usage of The Bethesda System (TBS) criteria.
33 34		Stability
35 36 37 38 39 40 41 42		Provide real-time studies from three different manufacturing lots which include data to demonstrate the stability of any reagents and/or buffers under expected shipping, handling, and storage conditions including various extremes of temperature, humidity, and light. This includes pre- collection buffer solutions and post-collection preservation of cellular morphology.

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1	C.	Clinical	Studies	
2		1 1	Stades Destan	
э Л		1	Study Design	
5		1	Provide the fo	allowing study design information:
6				nowing blady design miorination.
7			a. Patient	inclusion and exclusion criteria.
8				
9			Accou	nt for all patients and provide explanation for all patients
10			who ar	re excluded from study.
11				·
12		ł	b. Specifi	ic patient selection sampling plan.
13				
14			For me	ethods not limited to the conventional Pap smear:
15				
16			*	Prospective patient sampling is preferred so that within a
17				defined time frame all patients are consecutively entered
18				into the study. If not, provide the detailed statistical
19				sampling plan used. Frequently sponsors fail to use <u>all</u>
20				consecutive patients during the prespecified study time
21				period, therefore a clear description of the sampling
22				method must be provided. The method of sampling
23				should be selected to ensure the representativeness,
24				the target nonulation
25 26				the target population.
20			For me	thods limited to the conventional Pan smear
28				
29			*	Retrospective studies may be permitted for methods that
30				are solely based on the conventional Pap smear. e.g.
31				computer-assisted Pap test reading versus manual Pap test
32				reading. However, these will be considered a feasibility
33				study only.
34				
35				If negative archival slides are used, there must be an
36				IRB-approved written protocol detailing what will be
37				done with the study information from the new device. If
38				the new method yields abnormal diagnoses (including
39				ASCUS and above, AGCUS and above, etc.) on slides
40				that were previously diagnosed as negative, the laboratory
41				that diagnosed the conventional PAP slide must be
42				notified. The sponsor has the responsibility of

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1 2 3 4		monitoring and assuring appropriate patient follow-up. A confirmatory test must be performed according to TBS guidelines.
5	For all	devices:
6		
7	*	Select the endpoint of the clinical studies that will
8		support the clinical intended use.
9		
10		Novel and unproven devices may require the most
11		stringent data with endpoints that may include clinical
12		outcomes, i.e., effect of the device on morbidity and/or
13		mortality, colposcopy results, and directed biopsy results
14		as clinically indicated. Surrogate endpoints include
15		colposcopy with directed biopsy, conventional Pap test
10		versus thin-layer or monolayer Pap test, manually read
17		and interpreted Pap test vs. computer-assisted or
18		automated Pap test.
19	*	Define study period from which the notionts will be
20	Ŧ	related
21		selecicu.
22	*	Dravida the type and number of study contars
25	<b>.</b>	Provide the type and number of study centers.
24		Carefully select clinical study sites to represent the
25		spectrum of demographic features appropriate for the
20		study of certical pathology and include low prevalence
28		and high prevalence populations for the diseases for
20		which the device is to be used
30		which the device is to be used.
31	*	Provide the number of natients per center
32		riovide the number of putonts per conten.
33		Study adequate numbers of patients with confirmed
34		positive and confirmed negative results from each site to
35		allow for site stand-alone analysis and to support the
36		difference detected in clinical sensitivity and clinical
37		specificity at the pre-specified statistical significance level
38		and power.
39		•
40	*	Justify statistically all pooling of data from multi-center
41		studies. Collect sufficiently large numbers of specimens
42		

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1 2		from each study site to allow for stand-alone analysis at each site
3		cach site.
4	C.	Hypothesis testing
5	•••	xypoutone toking
6		Null and alternative hypotheses for 3 by 3 categorical
7		classification of ordinal data.
8		
9		The Bethesda System (TBS) cervical cytology diagnoses
10		represent qualitative or categorical data, e.g., negative, atvpical
11		squamous cells of undetermined significance (ASCUS), low
12		grade squamous intraepithelial lesion (LGSIL), high grade
13		squamous intraepithelial lesion (HGSIL), etc. TBS diagnoses
14		can be grouped into three treatment categories: negative, -
15		ASCUS, and LGSIL and above. These data are ordinally scaled
16		and may be formed into two (new device and reference)
17		stochastically ordered distributions.
18		·
19		Clearly define the null and alternative hypotheses to be tested for
20		the 3 by 3 ordinal classification tables as shown above. The null
21		hypothesis should be reasonably and consistently constructed
22		from the data analyses of the pilot studies for the reference and
23		the new device. The alternative hypothesis is the hypothesis
24		designed to include all possibilities not included in the null
25		hypothesis.
26		
27		For example:
28		
29		Null hypothesis = The two marginal distributions of clinical
30		outcomes (negative, ASCUS, LGSIL and higher) are equal
31		between the new and reference devices.
32		
33		Alternative hypothesis = The two marginal distributions of
34		clinical outcomes are not equal between the new and the
35		reference devices (two-sided). This hypothesis considers the
36		possibility that the device can be either better or worse than
37		the reference device.
38		
39		Perform appropriate statistical significance testing for the 3 by 3
40		ordinal classification data.
41		
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 If the null hypothesis is rejected in the above test for the 3 by 3 classification data, perform statistical significance tests for two 2 by 2 classification tables by combining ASCUS into the negative or positive (LGSIL and above) groups.

For each of the 2 by 2 tables, if the patient true disease status is known, calculate the clinical sensitivity values for the disease positive group and the clinical specificity values for the disease negative group for both the reference and new device. If the patient true disease status is not known, calculate the proportions of disease (LGSIL and higher) for the reference and new device respectively.

When the patient's true disease status is known, perform statistical significance testing for comparing two true clinical sensitivities and two true clinical specificities (or for two true proportions of disease if patient disease is not known) between the reference and new device. The appropriate procedure for the multiple comparison problem needs to be considered since each of the above 2 by 2 tables were reconstructed from the SAME 3 by 3 table when the 3 by 3 ordinal data leads to the rejection of the null hypothesis.

## d. Confidence Intervals

For the above 2 x 2 tables provide the 95 percent confidence intervals for true clinical sensitivity based on the diseasepositive confirmed group and clinical specificity based on the disease-free confirmed group for the new device if the patient disease status is known.

Also, provide the 95 percent confidence intervals for the differences of the two true clinical sensitivities and two true clinical specificities between the new and reference devices. This calculation can be useful in making clinical decisions.

When the patient's true disease state is not known, provide the true proportions of disease states, LGSIL or higher, along with the 95 percent confidence intervals for the difference between the two true proportions.

	Points to Consider: Cervical Cytol	ogy Devices, Version 7/25/94
1	e.	Justification of Sample Size
3		The following must be clinically and statistically pre-specified
4		for estimating the required sample size:
5		tor obtaintaing and required bumple size.
6		* Type Lerror (probability of rejecting a true null
7		hypothesis)
8		-,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
9		* Power (probability of rejecting a false null hypothesis)
10		
11		* Hypothetical clinical sensitivities and specificities (true
12		disease status is known)
13		······································
14		* Hypothetical proportions of LGSIL, HGSIL, and
15		carcinomas (true disease status is not known)
16		
17		* Disease prevalence _
18		
19		Collect a sufficient number of disease positive and negative
20		samples to test the difference to be detected in clinical
21		performance characteristics between the new device and the
22		standard reference device at the prespecified statistical
23		significance level and power.
24		
25	f.	Specify Statistical Analysis for ORDINAL Data for the
26		following parameters:
27		
28		Clinical Sensitivity
29		
30		Clinical Specificity
31		
32		Positive and Negative Predictive Values
33		
34		Inter- and Intra-observer agreement (masked)
35		
36	g.	The study population must simulate the intended population for
37		the device.
38		
39		* Sample and perform testing at 3-5 different geographical
40		sites. Selection of U.S. sites is strongly encouraged.
41		Unity one testing site may be the manufacturer's own test
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1 2			site. Simulate the testing and the use of the device for the site(s) for which the device is intended.
3			
4		*	Collect specimens from populations with a range of low
5			and high prevalences for the disease(s) to be detected by
6			the device according to its intended use.
7			C
8		*	Sample examples of all the cervical diseases and
9			conditions covered by the Bethesda System (TBS).
10			Clearly list in the intended use statement any exclusions
11			of diagnostic categories, e.g., "this device is not intended
12			to process, detect, interpret inflammatory cells, etc." or
13			"no examples of X condition were detected or analyzed
14			by this device during the premarketing studies. The
15			performance of this device is not known for X condition."
16			(X is a rare condition not readily detected in prospective
17			clinical studies of a size with sufficient power for most
18			conditions.)
19			
20			Design preclinical tests to support, at least theoretically,
21			that the device is safe and effective for all intended
22			conditions claimed, even the most rare ones.
23			
24			Design post-market studies to validate the safety and
25			effectiveness of the device for rare conditions using
26			sufficient numbers of clinical samples.
27			-
28		*	Plan for any other sampling requirement for other
29			conditions claimed in the intended use statement, e.g.,
30			processing of the samples for viral culture, nucleotide
31			studies, etc.
32			
33	2.	End-point: Cli	inical accuracy.
34		•	•
35		A validation of	of an in vitro diagnostic method requires an independent
36		reference end	point as a measure of accuracy (truth). Also, once a
37		reference met	hod is chosen, all results must be compared with this
38		method.	
39			
40		Measure how	well the device represents the true clinical condition of
41		the patient fro	om whom the sample was taken. Collect and analyze all
42		the results fro	m the screening cytotechnologist for all specimens and

	Points to Consider: Cervical Cytolo	gy Devices, Version 7/25/94
1	separa	tely analyze the results for all the specimens referred to the
2	pathol	ogist. Ensure that the protocol's criteria for referral from the
3	cytoted	chnologist to the pathologist are followed for all cases, both the
4	new de	evice and the reference method. Mask all comparisons of the new
5	device	to the reference method.
6		
7	The fo	llowing endpoints are listed in descending order of their power
8	as scie	ntific evidence for validation of a new device, beginning with the
9	gold st	tandard method:
10	-	
11	а.	Clinical outcome
12		
13		This is the gold standard reference for accuracy and clinical
14		utility but has the disadvantage that long term follow-up
15		intervals are required to evaluate gynecology cytology.
16		
17		Longer clinical studies may be required for devices that
18		are not based on conventional Pap smears because it may
19		not be possible to use archival cases for longitudinal
20		studies.
21		
22	b.	Cervical biopsy
23		
24		This method is less than a gold standard because of difficulty in
25		biopsy sampling. Biopsies may yield false negative results from
26		sampling error. Some of these patients may have a negative
27		cytology but actually have a pre-neoplastic or neoplastic
28		condition. Also, there are additional problems in that it is not
29		practical to biopsy a large number of patients with negative or
30		ASCUS or AGCUS cytology.
31		
32	с.	Colposcopy examination and directed cytology and/or biopsy
33		sampling
34		
35		If a colposcopy cervical cytology sample is to be used for
36		confirmation, ensure that a sufficient time has elapsed from any
37		previous cytology sampling to allow for cervical epithelium to
38		regenerate (4 to 6 weeks).
39		
40	3. Surrog	ate end-point: Specimen accuracy.
41	-	
42		A conventional Pap smear with refereed manual microscopic

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1 2 3			reading may be used as a data subset to resolve discrepancies and to assess specimen accuracy.
ј Л			Marked manual microscopy is an accortable sume sate reference
4			masked manual microscopy is an acceptable surrogate reference
5			conventional Pan smoor and as a comparison method for thin
0 7			laver or monolayer slide preparations. Record the masked
8			referred diagnoses before performing any unmasked comparison
0 Q			or "consensus" readings of slides or images
10			or consensus readings of sinces of images.
10			Mask each cytotechnologist and nathologist to the results of the
12			comparison method and each other's results for the conventional
13			Pan smear and for the new device e.g. an automated or semi-
14			automated image analysis or computer-assisted cell locator
15			device
16			GU 1200.
17			Document that the masking code will not be broken until all the
18			data is gathered and recorded. It is important to maintain the
19			masking code until the final data analysis.
20			
21	3.	Sampl	e collection
22			
23		a.	Collection device for cervical specimens.
24			•
25			Limit cervical sampling collection devices to those that are FDA
26			cleared. The device must sample the endocervix and the
27			exocervix. Data should be collected to ensure TBS criteria for
28			adequate cellularity, endocervical component, etc. The Ayres
29			spatula used alone is not sufficient for endocervical sampling.
30			The Brush used alone is not sufficient for exocervical sampling.
31			
			A combination of collection devices or a combination device,
32			A combination of collection devices or a combination device, e.g., cervical broom provide sufficient sampling of the
32 33			A combination of collection devices or a combination device, e.g., cervical broom provide sufficient sampling of the endocervix and exocervix.
32 33 34			A combination of collection devices or a combination device, e.g., cervical broom provide sufficient sampling of the endocervix and exocervix.
32 33 34 35			A combination of collection devices or a combination device, e.g., cervical broom provide sufficient sampling of the endocervix and exocervix. One disadvantage is that some brushes may cause
32 33 34 35 36			A combination of collection devices or a combination device, e.g., cervical broom provide sufficient sampling of the endocervix and exocervix. One disadvantage is that some brushes may cause excessive bleeding and may not be used during
32 33 34 35 36 37			A combination of collection devices or a combination device, e.g., cervical broom provide sufficient sampling of the endocervix and exocervix. One disadvantage is that some brushes may cause excessive bleeding and may not be used during pregnancy.
32 33 34 35 36 37 38	·		A combination of collection devices or a combination device, e.g., cervical broom provide sufficient sampling of the endocervix and exocervix. One disadvantage is that some brushes may cause excessive bleeding and may not be used during pregnancy.
32 33 34 35 36 37 38 39		b.	A combination of collection devices or a combination device, e.g., cervical broom provide sufficient sampling of the endocervix and exocervix. One disadvantage is that some brushes may cause excessive bleeding and may not be used during pregnancy. Devices that are not based on collection, preparation, and
32 33 34 35 36 37 38 39 40	·	ь.	<ul> <li>A combination of collection devices or a combination device, e.g., cervical broom provide sufficient sampling of the endocervix and exocervix.</li> <li>One disadvantage is that some brushes may cause excessive bleeding and may not be used during pregnancy.</li> <li>Devices that are not based on collection, preparation, and observation of conventional Pap smear slides, e.g., thin or</li> </ul>
32 33 34 35 36 37 38 39 40 41		b.	<ul> <li>A combination of collection devices or a combination device, e.g., cervical broom provide sufficient sampling of the endocervix and exocervix.</li> <li>One disadvantage is that some brushes may cause excessive bleeding and may not be used during pregnancy.</li> <li>Devices that are not based on collection, preparation, and observation of conventional Pap smear slides, e.g., thin or monolayer slides.</li> </ul>

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1 2 3 4 5		An Institutional Review Board (IRB) approval and informed consent is required if a patient is to be screened by a method that does not allow for a conventional Pap test (conventional Pap smear read manually by a cytotechnologist with potential for referral to a pathologist).
7 8		Specimen collection may be handled in one of the following two ways:
9		
10		If a Pap smear is to be made, collect the cervical sample and
11		prepare a conventional Pap smear in the usual fashion before
12		performing any of the steps for the non-Pap smear slide. The
13		sample for the new methodology, e.g., this layer or monolayer
14		suspension may be made from the residual material on the
15		conection spatula, orush or oroom.
10		If no conventional Dan amoun is made before compling for the
19		this layer or monolever preparation, randomize the study
10		subjects into two groups: One group will have a conventional
20		Pan smear and the other group will be sampled for the non-
20		conventional method e g thin layer or monolayer slides
21 22		Appropriate statistical design is needed to ensure equal or nearly
23		equal numbers of samples in each of the two groups. This
25		method may require an Investigational Device Exemption (IDE)
25		and signed patient consent forms as well as an IRB approval if
26		the study is a PMA.
20		
28		The end-point for these studies is a comparison of the proportion
29		of TBS diagnoses found in each of the two study groups.
30		
31		See attached addendum for additional sampling considerations.
32		r
33	4.	Sample processing
34		
35		Consideration must be given to the method for processing the cervical
36		sample. The information and data submitted for a smear would vary
37		from that of a suspension or a differential separation of the cervical
38		sample. The differential separation of cervical cells from mucus, red
39		blood cells, leukocytes, and acellular material would require proof that
40		diagnostic and contextual cells were not lost in the processing of the
41		sample. The method of separation, filtration or centrifugation, must be
42		addressed as well as the type of fixation.

	Points	to Consid	er: Cervical Cytology Devices, Version 7/25/94
1			5. Slide preparation
2 3 4 5 6 7			Issues involving the transfer of the cervical specimen to the slide will vary according to the method used. The transfer of cells from a suspension by filtration, centrifugation, or sedimentation is vastly different from a direct smear.
8			6. Cell finding-locating device
9 10 11 12 13 14 15			Issues to be addressed for this type of device include the theory of operation, the sensitivity and specificity of the image processing for the selection of cells. Does the device provide for marking of suspicious cells? What is the operator/instrument interaction? What type of long-term record is provided? Documentation for software that was designed as part of the cell-locating device should be provided.
17			7. Image interpretation
18 19 20 21 22 23 24 25			For devices that provide an image interpretation, issues to be addressed include the theory of operation, the comparison with conventional microscopy, operator instrument interaction, long-term record keeping, and a hazard analysis and software documentation conforming to the criteria outlined in <i>Reviewer Guidance For Computer Controlled Medical Devices Undergoing</i> $510(k)$ Review.
26 27	Ш.	Data	
28 29 30 31		А.	Record keeping Stratify data and results by TBS diagnoses for claimed intended use by screening cytotechnologist and pathologist. The following terminology should
32 33 34 35 36 37			be used: high grade squamous intraepithelial lesion (HGSIL), low grade squamous intraepithelial lesion (LGSIL), atypical squamous cells of undetermined significance (ASCUS), atypical glandular cellular of undetermined significance (AGCUS), etc. The definition of ASCUS should be confined to the TBS category. The descriptive diagnoses pertaining to presence of inflammation and infection as well as reactive changes and specimen
38 39			adequacy should also be noted.
40 41		В.	Data integrity
42			1. Retain all original work sheets and make them available for planned or

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1	unannounced FDA inspections. In cases where the original form does
2	not remain with the study, all transcribed data should show the initials
3	of the recorder and the date of transcription. If original work sheets
4	may not be available at a later date, it is important to provide a
5	verification for the transcribed data which should include the initials of
6	the individual verifying the transcription and the date of verification.
7	
8	Document in writing any correction or modification of original work
9	sheets. Corrections or modifications should follow standard good
10	laboratory practices in that a single line is drawn through the potation
11	that is changed. The correction is written above the line. Record the
12	date and the initials of individual making any changes
13	date and me manuals of mervidual maxing any changes.
14	Ensure the integrity of data on computerized work sheets (databases) so
15	that there is permanent recording of date of input of all data and all
16	that affect is permanent recording of date of input of an data and an
17	-
19	Mask reading interpretation and recording of data. Mask all data
10	mask reading, interpretation, and recording of data. Mask an data
20	analysis and interpretation.
20	2 At the intended site of use conventional manually prepared Pan slides
21	2. At the included site of use, conventional manually prepared r ap sinces
22	and sides prepared with a new device of read by a new device, e.g.,
23 24	diamages that fall between TPS actagories. It is most likely that these
24	alides will be treated like the discretion of the higher rather than the
25 26	sinces will be dealed like the diagnosis of the higher father than the
20	lower 1 by diagnosis bracketing the provisional diagnosis. The side
21	would be referred by the cytolechnologist to the next higher trained
28	odserver.
29	If the methole sight discussion falls a subscrabble hadman from TDC
30	If the pathologist's diagnosis fails equivocally between two TBS
31	diagnostic categories, it would be expected that the pathologist would
32	refer the slide to another pathologist or would note the equivocal
33	diagnosis in the report to the clinician for appropriate follow-up.
34	
35	Record all diagnoses on the work sheet that are not definite 1BS
36	diagnoses, e.g., hedging diagnoses such as tentative, provisional,
37	presumptive, rule out, R/O, or question mark ?, as the next higher
38	grade. Establish a written algorithm in the protocol for all possible
39	diagnostic variations before the study begins.
40	
41	Ensure that data entry personnel are aware of TBS diagnostic criteria.
1	

	Points	to Consider: Cervic	al Cytology Devices, Version 7/25/94
1			reviewer in the cytology protocol, i.e., all slides that are not
2			unequivocany negative.
2		2	If the device is commuter excisted world, shownal calls by mound
-+ 5		Ј.	in the device is computer-assisted, verify abnormal cells by manual
5			meroscopy to check and re-check cens identified as abnormal.
7			If it is not possible to check abnormal cells by manual microscopy
8			iustify the safety and effectiveness of the new methodology
9			Justify the survey and effectiveness of the new methodology.
10		4.	If the device displays abnormal cells on a computer monitor, document
11			how these images are stored for additional observers to re-analyze.
12			
13		5.	Provide instructions in the study protocol to ensure adherence to the
14			manufacturer's protocol by the cytotechnologists and pathologists.
15			
16		6.	Devices that allow for little or no opportunity for human intervention, in
17			particular a software-controlled "black box-type" devices, may require
18			software validation studies to document that the device can be expected
19			to maintain reasonable performance for its intended purpose with the
20			full range of expected specimens.
21			
22	IV.	<b>Training Req</b>	uirements
23			
24		State the kind	and amount of training requirements for any step and interpretation that
25		differs from t	ne conventional Pap test (manually prepared smear, manually processed,
26		and read and	interpreted without computer assistance).
27		<b>D</b> 1.	
28		Recommend t	he appropriate procedures for the laboratorian and laboratory to convert
29		from convention	onal Pap test methodology to that of the new device.
30		Some	training issues to consider and
37		Some	d'anning issues to consider ale.
33		Will H	he new device affect the appearance of the finished slide?
34		•• III U	to now device after the appearance of the mission side?
35		Will th	e slide be a conventional Pan smear or a thin layer or monolayer slide?
36			
37		Will th	e new slide methodology contain all of the components of the
38		conver	ntional slide?
39			
40			If not, what is the possibility of misdiagnoses from loss of that (those)
41			components?
42			

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1		How will these issues concern the clinician during sample collection, sample
2		preparation, slide preparation, and final interpretation of result?
3		How will there issues affect the suitstached said during sample processing
4 5		slide preparation identification of abnormal cells, and interpretation of cells
6		from the preparation?
7		mont de propulation.
8		Consider how these issues affect the pathologist in identification of abnormal
9		cells and interpretation of abnormal cells?
10		•
11	V.	Workload Limit
12		
13		All gynecologic cytology devices submitted for FDA approval or clearance must have
14		an evaluation of their workload limit. This applies to devices for making cell
15		suspensions for thin layer slides, computer-assisted cell locator devices, semi-
16		automated and automated computer-assisted image analyzers, and any other device
1/		used in the preparation, reading, and interpretation of gynecologic cytology specimens.
10		Data must be provided to assess and evaluate the rangue factor in order to establish
19 20		appropriate workload mints.
21	vī	References
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