Department of Health and Human Services Food and Drug Administration

STANDARDS DATA REPORT FOR 510(k)s

(To be filled in by applicant)

This report and the Summary Report Table are to be completed by the applicant when submitting a 510(k) that references a national or international standard. A separate report is required for each standard referenced in the 510(k).

TYPE OF 510(K) SUBMISSION		
Traditional Special Abbreviated STANDARD TITLE ¹		
STANDARD TITLE		
Please answer the following questions	Yes	No
Is this standard recognized by FDA ² ?		
FDA Recognition number ³	#	
Was a third party laboratory responsible for testing conformity of the device to this standard identified in the 510(k)?		
Is a summary report ⁴ describing the extent of conformance of the standard used included in the 510(k)?		
Does the test data for this device demonstrate conformity to the requirements of this standard as it pertains to this device?		
Does this standard include acceptance criteria? If no, include the results of testing in the 510(k).		
Does this standard include more than one option or selection of tests?		
Were there any deviations or adaptations made in the use of the standard? If yes, were deviations in accordance with the FDA supplemental information sheet (SIS) ⁵ ?		
Were deviations or adaptations made beyond what is specified in the FDA SIS?		
Were there any exclusions from the standard?		
Is there an FDA guidance ⁶ that is associated with this standard?		
Title of guidance:		
¹ The formatting convention for the title is: [SDO] [numeric identifier] certification body involved in conformance assessment	ent to this	

- ¹ The formatting convention for the title is: [SDO] [numeric identifie [title of standard] [date of publication]
- ² Authority [21 U.S.C. 360d], www.fda.gov/cdrh/stdsprog.html
- 3 http://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfStandards/ search.cfm
- ⁴ The summary report should include: any adaptations used to adapt to the device under review (for example, alternative test methods); choices made when options or a selection of methods are described; deviations from the standard; requirements not applicable to the device; and the name and address of the test laboratory or
- certification body involved in conformance assessment to this standard. The summary report includes information on all standards utilized during the development of the device.
- ⁵ The supplemental information sheet (SIS) is additional information which is necessary before FDA recognizes the standard. Found at http://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfStandards/ search.cfm
- 6 The online search for CDRH Guidance Documents can be found at www.fda.gov/cdrh/guidance.html

EXTENT OF STANDARD CONFORMANCE SUMMARY REPORT TABLE STANDARD TITLE **CONFORMANCE WITH STANDARD SECTIONS*** SECTION TITLE SECTION NUMBER CONFORMANCE? Yes No N/A TYPE OF DEVIATION OR OPTION SELECTED * DESCRIPTION JUSTIFICATION SECTION NUMBER SECTION TITLE CONFORMANCE? Yes No N/A TYPE OF DEVIATION OR OPTION SELECTED * DESCRIPTION JUSTIFICATION SECTION TITLE SECTION NUMBER CONFORMANCE? Yes N/A No TYPE OF DEVIATION OR OPTION SELECTED * DESCRIPTION JUSTIFICATION * For completeness list all sections of the standard and indicate whether conformance is met. If a section is not applicable (N/A) an explanation is needed under "justification." Some standards include options, so similar to deviations, the option chosen needs to be described and adequately justified as appropriate for the subject device. Explanation of all deviations or description of options selected when following a standard is required under "type of deviation or option selected," "description" and "justification" on the report. More than one page may be necessary. Types of deviations can include an exclusion of a section in the standard, a deviation brought out by the FDA supplemental information sheet (SIS), a deviation to adapt the standard to the device, or any adaptation of a section. **Paperwork Reduction Act Statement**

Public reporting burden for this collection of information is estimated to average 1 hour per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to:

Department of Health and Human Services Food and Drug Administration Office of Chief Information Officer 1350 Piccard Drive, Room 400 Rockville, MD 20850

An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB control number.