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

## PREMARKET NOTIFICATION [510(K)] STATUS REQUEST AND RESPONSE

**Single Form** – To be used for both your request and FDA's response. Requesters should fill in the information on the top half of this form (Request Section) and fax (or mail) this form to the FDA at the fax # or address listed below. The FDA will complete the information on the bottom half (Response Section) and return by fax (or mail).

REQUEST SECTION (To be completed by requester)						
From (To)						
REQUESTER NAME						
MAILING ADDRESS						
FAX NUMBER						
REQUESTER'S AFFILIATION WITH THE SUBMITTER OF THE 510(K)						

**Requester Certification:** I certify that I am an authorized representative of the submitter of the following 510(k) and that all information provided herein is truthful to the best of my knowledge. Please provide me with information related to the status of the following 510(k) submission via *(mark one)*: FAX  $\Box$  or MAIL  $\Box$ 

510(K) NUMBER	REQUESTER SIGNATURE
SPONSOR'S NAME AND ADDRESS	PRODUCT NAME
	DATE LOGGED IN BY FDA (ODE) – as identified in acknowledgement letter

## **RESPONSE SECTION** (To be completed by FDA)

NOTE: THIS DOCUMENT IS INTENDED ONLY FOR THE USE OF THE PARTY TO WHOM IT IS ADDRESSED AND CONTAINS INFORMATION THAT IS PRIVILEGED, CONFIDENTIAL, AND PROTECTED FROM DIS-CLOSURE UNDER APPLICABLE LAW.

If you are not the addressee, or authorized to deliver this document to the addressee, you are hereby notified that review, disclosure, dissemination, copying, or other action based on the content of this communication is not authorized. If document has been received in error, please notify FDA by phone and return via mail.

Reviewing Branch:	. LA	ST ACTION	AND DATE
Please be advised that the average total time (time for FDA review plus	6		
time spent awaiting any additional data) for review of a device assigned to			
this branch has been days over the last 6 months.			

Place in Queue: You	ur 510(k) has bee	en assigned to a	reviewer a	ind is #	in line for	r that reviewer to work
on. The length of time	e that it will take	for the reviewer	to get to y	our 510(k) a	and to review it	will depend on many
factors, such as the	complexity of the	e 510(k)'s that	are in line	ahead of y	ou, and other	work assigned to the

reviewer, for example the review of investigational device exemption submissions. Due to these variables, we cannot estimate a completion date for review of your 510(k). However, future inquiries can give you an idea of how your 510(k) is progressing.

FDA RESPONSE DATE

Please do not request another status report prior to 30 days from the FDA response date.

Public Reporting burden for this collection of information is estimated to average 0.2 hour per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing 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:

Food and Drug Administration CDRH (HFZ-220) 2094 Gaither Road Rockville, MD 20850