DEPARTMENT OF HEALTH AND HUMAN SERVICES FOOD AND DRUG ADMINISTRATION

CORH PREMARKET REVIEW SURMISSION COVER SHEET

Form Approval
OMB No. 9010-0120
Expiration Date: August 31, 2010.

		DIVIDUOIV (JOVEN SHI		See OMB Sta		
Date of Submission	User Fee Payment		FDA Submissi	on Document	Numbe	r (if known)	
SECTION A		TYPE OF S	UBMISSION				
PMA	PMA & HDE Supplement	PD		510(k)			Meeting
Original Submission Premarket Report Modular Submission Amendment Report Report Amendment Licensing Agreement	Regular (180 day) Special Panel Track (PMA Only) 30-day Supplement 30-day Notice 135-day Supplement Real-time Review Amendment to PMA & HDE Supplement Other	Original PI Notice of C Amendme	Completion	Original Submit Traditional Special Abbreviated section I, Par Additional Infor	(Complete ge 5)	Pre- Pre- Day Agro	-510(K) Meeting -IDE Meeting -PMA Meeting -PDP Meeting -100 Meeting eement Meeting ermination Meeting er (specify):
IDE	Humanitarian Device Exemption (HDE)	Class II Exemp	otion Petition	Evaluation of Aut	nation	Othe	er Submission
Original Submission Amendment Supplement	Original Submission Amendment Supplement Report Report Amendment	Original St	ubmission Information	(De Novo) ssion	5136 Othe	· ·· ·
Have you used or cited Stand	dards in your submission?	Yes No	(If Yes,	please complete Se	ction I, Page 5	 5)	
SECTION B	SUBM	IITTER, APPLI	CANT OR SP	ONSOR			
Company / Institution Name			Establishment	Registration Number	(if known)		
Division Name (if applicable) Phone Number (including area code) ()							
Street Address	FAX Number (i	ncluding area code)					
City	State / Province	/ Province ZIP/Postal Code Country					
Contact Name							
Contact Title			Contact E-mail	Address			
SECTION C	APPLICATION CORRES	PONDENT (e.	g., consultan	t, if different from	above)		
Company / Institution Name							
Division Name (if applicable)			Phone Number	(including area code)		
Street Address			FAX Number (i	ncluding area code)			
			()	<u> </u>			
City	State / Province ZIP/Postal Code Cou			Country			
Contact Name							
Contact Name							
Contact Title			Contact E-mail	Address			

SECTION D1 REA	SON FOR APPLICATION - PMA, PDP, OR H	IDE
Withdrawal Additional or Expanded Indications Request for Extension Post-approval Study Protocol Request for Applicant Hold Request for Removal of Applicant Hold Request to Remove or Add Manufacturing Site Process change: Manufacturing Sterilization Packaging Other (specify below) Response to FDA correspondence:	Change in design, component, or specification: Software / Hardware Color Additive Material Specifications Other (specify below) Labeling change: Indications Instructions Performance Shelf Life Trade Name Other (specify below)	□ Location change: □ Manufacturer □ Sterilizer □ Packager □ Report Submission: □ Annual or Periodic □ Post-approval Study □ Adverse Reaction □ Device Defect □ Amendment □ Change in Ownership □ Change of Applicant Address
Other Reason (specify):		
SECTION D2	REASON FOR APPLICATION - IDE	
New Device New Indication Addition of Institution Expansion / Extension of Study IRB Certification Termination of Study Withdrawal of Application Unanticipated Adverse Effect Notification of Emergency Use Compassionate Use Request Treatment IDE Continued Access	Change in: Correspondent / Applicant Design / Device Informed Consent Manufacturer Manufacturing Process Protocol - Feasibility Protocol - Other Sponsor Report submission: Current Investigator Annual Progress Report Site Waiver Report	Repose to FDA Letter Concerning: Conditional Approval Deemed Approved Deficient Final Report Deficient Progress Report Deficient Investigator Report Disapproval Request Extension of Time to Respond to FDA Request Hearing Request Hearing
Other Reason (specify):		
SECTION D3	REASON FOR SUBMISSION - 510(k)	
New Device	Additional or Expanded Indications	Change in Technology
Other Reason (specify):		

FORM FDA 3514 (9/07) PAGE 2 OF 5 PAGES

SECTION E		which substar	ADDITION ntial equivalence		ORMATION	ON 51	0(K) SUB	MISS	NOI		or statement concerning,
1	or devices to v	2	iliai equivalence		3 4					safety and effe	ectiveness information	
5		6		7				• • • • • • • • • • • • • • • • • • •		(k) summary attached (k) statement		
	devices to wh		al equivalence is		(if known)							
illioilliation on	510(k) N		ai equivalence is		ade or Proprieta	arv or M	odel i	 Name				nufacturer
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1			1							1		
2			2							2		
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SECTION F	.	- 25	RODUCT INFO		ON APPL	CATIO	N E-2	O AL-1	A.D.E		ATIONS	
	sual name or cl		IODOOT IIII	ZI IIW/AI I	ON AITEN	OATIO		<i>5 1</i> (1)			ATIONS	
Trade or F	Proprietary or M	lodel Name for	or This Device						Мо	del N	lumber	
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5									5			
FDA documen			d submissions (I	egardless	of outcome)							
1		2	3			4				5		6
7		8	9			10				11		12
Data Included	in Submission	<u></u> □ ι	Laboratory Testir	ng	☐ Aı	nimal Tr	als				Human Trials	
SECTION G			DDUCT CLAS	SIFICA	TION - APPL	ICATI	ON T				CATIONS	
Product Code	C.F.	.R. Section (i	if applicable)					Device	e Class	S		
Classification	Panal							ļ 🗆 (Class I		Class II	
Classification Panel						Class III Unclassified						
Indications (fro	om labeling)											

FORM FDA 3514 (9/07) PAGE 3 OF 5 PAGES

Note: Submission of this or 2891a Device Establish	information does not affect the ne ment Registration form.	eed to submit a 2891	FDA Document Number (if known)					
SECTION H Original Add Delete Company / Institution Name	TERILIZATION SITES RELATING TO A SUBMISSION Manufacturer Contract Sterilizer Contract Manufacturer Repackager / Relabeler Establishment Registration Number							
Division Name (if applicated applicated street Address	Phone Number (including area code) () FAX Number (including area code) ()							
City			State / Province		ZIP/Postal Code	Country		
Contact Name		Contact Title	Contact E-mail Address					
Original Add Delete Company / Institution Name			Manufacturer Contract Sterilizer Contract Manufacturer Repackager / Relabeler Establishment Registration Number					
Division Name (if applicate	Phone Number (including area code) ()							
Street Address City	FAX Number (including area code) () State / Province ZIP/Postal Code Country							
Contact Name Contact Title			Contact E-mail Address					
Original Add Delete Company / Institution Name			☐ Manufacturer ☐ Contract Sterilizer ☐ Contract Manufacturer ☐ Repackager / Relabeler Establishment Registration Number					
Division Name (if applicated applicated street Address	Phone Number (including area code) () FAX Number (including area code)							
City			() State / Province ZIP/Postal Code Country					
Contact Name Contact Title					Contact E-mail Addr	ess		

FORM FDA 3514 (9/07) PAGE 4 OF 5 PAGES

SECTION I UTILIZATION OF STANDARDS									
Note: Complete this section if your application or submission cites standards or includes a "Declaration of Conformity to a Recognized Standard" statement.									
1	Standards No.	Standards Organization	Standards Title	Version	Date				
2	Standards No.	Standards Organization	Standards Title	Version	Date				
3	Standards No.	Standards Organization	Standards Title	Version	Date				
4	Standards No.	Standards Organization	Standards Title	Version	Date				
5	Standards No.	Standards Organization	Standards Title	Version	Date				
6	Standards No.	Standards Organization	Standards Title	Version	Date				
7	Standards No.	Standards Organization	Standards Title	Version	Date				
	Please include any additional standards to be cited on a separate page.								

Public reporting burden for this collection of information is estimated to average 0.5 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-342) 9200 Corporate Blvd. Rockville, MD 20850

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FORM FDA 3514 (9/07) PAGE 5 OF 5 PAGES