

Institutional Review Board

INVESTIGATIONAL DEVICE FORM

The Christ Hospital Institutional Review Board SOP 2.01 Guidelines for Protocol Submission							
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	ı						
IRB #:							
PI Name:							
Study Title:							
Name of Do	dico:						
Name of Device: Model:							
Manufacture	-r:						
Wanaractar							
Has this devi	ice be	en approved by the FDA?	Yes	☐ No			
If yes, Does the proposed research use the device for the purposes for which it has been approved?			Yes	□No			
Indicate dev	ice st	atus (check as many as apply)					
	tiona	l Device Exemption (IDE) Number:					
or	tiona	r bevice exemption (ibe) Number					
FDA Exemption Letter							
Significant Risk Device							
Non-Significant Risk (NSR) Device							
-Attach a letter from the sponsor discussing the reasons for the risk classification.							
		6					
Commercially available							
Other	•						
Will device b	e bill	ed to participants?	Yes	☐ No			
If No, who w	vill pa	y for this device:					
Will a Data and Safety Monitoring Board (DSMB) review the data from this study?							
Ves /sere	nloto	Itoms a al					
Yes (complete Items a-e) No (and the research involves greater than minimal risk, complete Item f)							
a. Who will make up the DSMB:							

IDE Form

Version 3, IRB Review: 12/20/20



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b.	Will there be an Interim Analysis?:						
c.	At what points will the data be reviewed?:						
d.	Describe how the plan will work:						
e.	State any differences in the serious adverse event reporting between this particular proto FR 312 (or enter "None"):	ocol and 45 CF	R 46 or 21				
f.	Describe the plan for monitoring the data to ensure the safety of participants (Describe what data will be monitored, how often it will monitored, who will monitor it, what data will be evaluated, and the decisions that will be made in response to the evaluation):						
Report of handling, shipping and storage, release procedure and documentation of investigational devices (Note: Must maintain records for a period no less than 2 years after FDA approval or longer, if required by the sponsor, in accordance with FDA regulations)							
Do	es the investigator hold an IDE?	Yes	☐ No				
	Il device handling, storage, and documentation be tracked without an independent initor (e.g, CRO, other specified person)?	Yes	☐ No				
If No, name of monitor:							
List personnel who are trained and qualified to use the device:							
Describe the plan for handling, shipping and storage, control, and dispensing of the device so that only authorized investigators will use the device and they will use the device only in participants who have provided consent:							
Name (Printed or Typed)							
	natura	Data	Data				
ગાg	nature	Date					

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