DEPARTMENT OF HEALTH AND HUMAN SERVICES

FOOD AND DRUG ADMINISTRATION

CDRH Medical Device Reporting 10903 New Hampshire Avenue WO Bldg. 66, Room 3217 Silver Spring, MD 20993-0002 or email: MDRTHelpDesk@FDA.HHS.GOV

PART 1 INSTRUCTIONS

MEDICAL DEVICE REPORTING ANNUAL USER FACILITY REPORT

OMB: 0910-0437 Exp. Date: 3/31/2025

PART 1 - COVER SHEET

If MDR reports were not submitted to either the FDA or a device manufacturer during this reporting period, DO NOT submit an annual report.

Complete one copy of th above. This report shoul						
1. REPORT PERIOD			2. USER FA	ACILITY ID (CMS OF	R FDA PROVIDED NU	IMBER)
J	AN - DEC YYYYY			·		·
3. USER FACILITY INFORMATION	ON		4. USER F	ACILITY CONTACT	INFORMATION	
a. Name			a. Name)		
b. Street Address			b. Stree	t Address		
c. City	d. State	e. ZIP Code	c. City		d. State	e. ZIP Code
f. Country/Postal Code (if not	U.S.)			ry/Postal Code <i>(if no</i> hone Number <i>(Includ</i>)	t U.S.) de area code and exte	nsion)
5. TOTAL NUMBER OF REPOR	TS ATTACHED OR SUMM	MARIZED		, 		
a. Lowest Report Number	(CMS or FDA Provided N	(No.) (N		quence No.)		
b. Highest Report Number	(CMS or FDA Provided No	- <u>-</u> (Y		quence No.)		
For each report in the range of MedWatch FDA Form 3500A fo above range that are not include	r the event that was sent	to FDA and/or				
6. SIGNATURE OF CONTACT				7. DATE OF REPOR	RT	- y y y

This section applies only to the requirements of the Paperwork Reduction Act of 1995.

DO NOT SEND YOUR COMPLETED FORM TO THE PRA STAFF EMAIL ADDRESS BELOW.

The public reporting burden time for this collection of information is estimated to average 1 hour per response, including the time to review instructions, search existing data sources, gather and maintain the data needed and complete and review the collection of information. Send comments regarding this burden estimate or any other aspect of this information collection, including suggestions for reducing this burden, to:

Department of Health and Human Services Food and Drug Administration Office of Operations Paperwork Reduction Act (PRA) Staff PRAStaff@fda.hhs.gov 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.

MEDICAL DEVICE REPORTING ANNUAL USER FACILITY REPORT

PART 2 - SUMMARY OF EVENT

PART 2 INSTRUCTIONS

If photocopies of previously submitted FDA Form 3500A (MedWatch) are not provided for each MDR reportable event, complete one copy of the following for each MDR report submitted to FDA and/or the manufacturer during the calendar year covered by this Annual Report.

calendar year covered by this Annual Report.						
1. USER FACILITY EVENT REPORT NUMBER						
-	-					
(CMS or FDA Provided No.)	(Year) (Sequence No.)					
2. WHERE WAS REPORT SUBMITTED? (Check all that apply)						
FDA Manufacturer Distributor Other						
3. MANUFACTURER INFORMATION	4. DEVICE INFORMATION					
a. Name	a. Brand Name					
	b. Common Name					
h Olyant Address	_					
b. Street Address	c. Model Number					
	S. Model Hambel					
c. City d. State e. ZIP Code	d. Serial Number					
	e. Lot Number					
f. Country/Postal Code (if not U.S.)	1					
	f. Catalog Number					
5. BRIEF DESCRIPTION OF EVENT	Ш					