DEPARTMENT OF HEALTH AND HUMAN SERVICES FOOD AND DRUG ADMINISTRATION CDRH Medical Device Reporting	MEDI	MEDICAL DEVICE REPORTING ANNUAL USER FACILITY REPORT PART 1 - COVER SHEET			OMB: 0910-0437 Exp. Date: 8/31/2015	
P.O. Box 3002 Rockville, MD 20847-3002	Р					
If MDR reports were not submitted to either the FDA or a device manufacturer during this reporting period, DO NOT submit an annual report.						
PART 1 INSTRUCTIONS						
Complete one copy of the following inform	ation as a cove	r nade for	the annual report :	and return to th	address listed	
above. This report should NOT include rep						
1. REPORT PERIOD		2. USER FACILITY ID (HCFA OR FDA PROVIDED NUMBER)				
JAN - DEC $\overline{Y} \overline{Y} \overline{Y} \overline{Y} \overline{Y}$	-					
3. USER FACILITY INFORMATION			ACILITY CONTACT I			
a. Name		a. Nam				
b. Street Address		b. Street Address				
c. City d. State	e. ZIP Code	c. City		d. State	e. ZIP Code	
f. Country/Postal Code (if not U.S.)			f. Country/Postal Code (if not U.S.)			
			ohone Number (Include	a area code and e	stansion)	
)	died Loue and C.	(lension)	
5. TOTAL NUMBER OF REPORTS ATTACHED OR SU	UMMARIZED					
a. Lowest Report Number	(equence No.)			
b. Highest Report Number			equence No.)			
For each report in the range of report numbers liste MedWatch FDA Form 3500A for the event that was s above range that are not included in this report and exp	sent to FDA and/or					
6. SIGNATURE OF CONTACT			7. DATE OF REPOR	Т		
				//	<u> </u>	
This section applies of	and to the requireme	nts of the Pane	rwork Reduction Act of	1005		
***DO NOT SEND YOUR COM		-				
The public reporting burden time for this collection of search existing data sources, gather and maintain the of burden estimate or any other aspect of this information	lata needed and comp	plete and revie	w the collection of infor	mation. Send comm		
Department of Health and Human Food and Drug Administration Office of Chief Information Office Paperwork Reduction Act (PRA) S <i>PRAStaff@fda.hhs.gov</i>	er		gency may not conduct o ired to respond to, a colle displays a currently vali	ection of information	ı unless it	

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.

I. USER FACILITY EVENT REPORT NUMBER	
-	-
(HCFA or FDA Provided No.)	(Year) (Sequence No.)
2. WHERE WAS REPORT SUBMITTED? (Check all that apply)	
FDA Manufacturer Distributor Other	
B. MANUFACTURER INFORMATION	4. DEVICE INFORMATION
a. Name	a. Brand Name
	b. Common Name
b. Street Address	
D. Stieet Address	- Madel Mussher
	c. Model Number
c. City d. State e. ZIP Code	d. Serial Number
	e. Lot Number
f. Country/Postal Code (if not U.S.)	
	f. Catalog Number
5. BRIEF DESCRIPTION OF EVENT	