

U.S. NUCLEAR REGULATORY COMMISSION		Conversation Date: 5-20-05	
TELEPHONE CONVERSATION RECORD		Time: 11:15 am	
Mail Control or Report No.	License No.	Docket No.	
N/a	37-07722-04	030-03094	
Licensee/Applicant Participant(s):	Organization:	Telephone No.	
Mike Bieda, MP	Bryn Mawr Hospital	610-526-3372	
Person(s) Calling: Penny Lanzisera			
Subject: Part 21 Report (Event #41720)			
<p>Summary:</p> <p>Discussed with Mike the following:</p> <ol style="list-style-type: none"> 1) Event occurred on 5/17 (not 4/17). He will call operations center to correct. 2) Novoste device returned to manufacturer 5/18 3) Preliminary discussions with Novoste indicate that device was faulty, however, Novoste will contact Mike with additional information after their review. 4) Licensee did not use dual syringe system, however, he does not believe this was a contributing factor to the event, since it took an inordinate amount of pressure to get sources to return to the shield. After the event, he took the device and catheter containing the sources back to the hot lab and put a new syringe on the device. Even with the new syringe, he indicated that it took a lot of pressure to get the sources back in the shield. 5) The licensee treated the second location, because they believed the hang up of the sources during the treatment of the first location was caused by a kink in the catheter or a bad connection. They replaced the catheter and treated the second location, where the sources hung up again. 6) The licensee sent a faulty device report to the FDA. 			
Action Required/Taken: file			
Prepared By: Penny Lanzisera		Date: 5-20-05	