

Medtronic

Urgent Field Safety Notice

Pump Weld Update for HeartWare™ Ventricular Assist Device

(HVAD™)

Patient Management Recommendations

Model	Product Description	
1104	HVAD™ Pump Implant Kit	
MCS1705PU	HVAD™ Pump Implant Kit	

July 2022

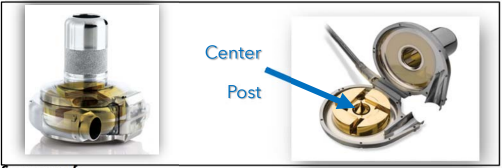
Medtronic Reference: FA1243 Phase III EU Manufacturer Single Registration Number (SRN): MF-000019976.

Dear Physician and Healthcare Professional:

Medtronic is providing this letter as a follow-up to our April 2022 communication titled "Urgent Medical Device Correction" (attached), which communicated a pump weld non-conformance with the HeartWare™ Ventricular Assist Device (HVAD™) System where (3) pumps were identified to have an impeller rotating non-concentrically and contacting the center post of the pump (see Figure 1: Pump Assembly). Medtronic's investigation was not able to conclusively isolate this issue to a specific subset of pumps. This communication provides updated information on the number of events, additional event details, and root cause.

There are no new patient management recommendations since the April 2022 communication. Consistent with the April communication, routine prophylactic explant of the HVAD device is not recommended, as risks associated with explantation may outweigh the potential benefits of continued HVAD therapy.

Figure 1: Pump Assembly



Summary of Complaints Confirmed

Since the April communication, Medtronic has received one (1) additional confirmed complaint related to this condition, for a total of four (4) complaints. In all four (4) complaints, each patient underwent a pump exchange due to suspicion of pump thrombus; however, no evidence of thrombus was found, and inspection of the explanted pumps revealed evidence of the impeller contacting the center post due to non-concentric rotation, consistent with corrosion of the center post magnet. Two (2) deaths were reported in association with these four (4) complaints: in one (1) case the patient underwent a cardiac transplant two (2) months after the pump exchange and died one month after the transplant; in the other case the patient died three (3) weeks after the VAD exchange.

Investigation of the four (4) complaints and bench testing of simulated corroded magnets indicates that an abnormal noise or vibration, commonly described as a "grinding" sound, was heard by either the patients or their clinicians. The grinding sound did not resolve after treatment for thrombus and is a likely initial indication that the impeller is contacting the center-post due to nonconcentric rotation. Over time, transient power spikes occurred on the logfiles causing [High Watt] alarms. This is unlike the gradual steady increase in power consumption typically seen with pump thrombus.

Although the root cause of the four complaints referenced above has been confirmed by analysis of the returned pumps, Medtronic continues to investigate complaints of suspected pump weld non-conformances.

Patients with affected devices may present with signs and symptoms that resemble pump

thrombus. It is not known if a patient's pump with this issue will present with the same signs/symptoms. In all four (4) instances the following were consistently encountered:

- Suspected Thrombus
- High Watt Alarms
- Grinding sound

The table below provides a summary of the four (4) confirmed complaints provided to Medtronic.

	Manufacturing Date	Implant Duration	Reported Signs and Symptoms
Complaint 1	Dec 2017	25 months	 Suspected thrombus
			 High watt alarms

Grinding sounds	
Vibration	
Reported Patient	Symptoms:
fatigue, light-hea	dedness and
dizziness, shortn	ess of breath
Complaint 2Jan 201828 months• Suspected throm	bus
Grinding sounds	
High watt alarms	
Elevated LDH lev	rel
Reported Patient	Symptoms: dark
urine	
Complaint 3May 201835 months• Suspected throm	bus
Grinding sounds	
High watt alarms	
Elevated LDH lev	rel
Reported Patient	Symptoms:
unknown	
Complaint 4April 201930 months• Suspected throm	bus
Grinding sounds	
High watt alarms	
Low flow alarms	
Elevated LDH lev	el
Reported Patient	Symptoms:
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Summary of Root Cause Investigation

As part of the investigation, Medtronic conducted a broad search of historical complaints and product returns to determine if there were additional pumps with suspected thrombus that may have a previously undetected corroded center-post magnet. As of July 2022, Medtronic completed an analysis of over 747 complaints from our returned product archives. Of those 747 complaints, Medtronic conducted a further analysis on 54 explanted pumps that had been returned to our analysis lab from 2012 to 2022. The returned pumps had allegations of thrombus, or a report of grinding sound or vibration. No additional occurrences of center-post magnet corrosion were found.

The investigation has determined that the occurrence of a weld crack is the result of a combination of factors that may include the initial presence of contamination in the weld area from previously applied substances as part of the manufacturing process, the misalignment of the cap and housing prior to welding, or the depth/thickness of the weld. Medtronic's investigation employed multiple methods to isolate the issue to a specific subset of pumps including quantifying weld thickness, alignment

indicators, and visual indications of contamination using over 8000 digital photos. Medtronic has found that there is not enough data available to conclusively isolate this issue to a specific subset of pumps.

Patient Management Recommendations

Consistent with the April 2022 communication, routine prophylactic explant of the HVAD device is not recommended, as risks associated with explantation may outweigh the potential benefits. Physicians should make the decision regarding explant and exchange of the HVAD pump on a case-by-case basis (is the patient a candidate for pump exchange, heart transplant, or pump explant for recovery), considering the patient's clinical condition and surgical risks. If a pump is explanted or exchanged for any reason, please return it to Medtronic for further analysis.

For patients presenting with any of the above signs and symptoms consider whether the clinical presentation could be due to a pump thrombus and treat accordingly. Please contact your Medtronic representative to provide details regarding the sequence of events and patient outcomes.

If patients present with these signs and symptoms listed above, please upload and submit all .csv logfiles to <u>https://autologs.medtronic.com</u>. Once on the website, please ensure to select the HVADlogs radio button and select "Urgent". Your Medtronic representative can assist with further logfile submission and analysis questions. Medtronic will analyze these logfiles and any other signs/symptoms as part of the ongoing investigation.

Customer Instructions

Medtronic records indicate that your site has patients that may still be on support; we request that you do the following:

- This notice must be shared with all those who need to be aware within your organization or any organization where patients have been transferred.
- Please complete the enclosed Customer Acknowledgement Form and email to your Medtronic Representative.

Additional information

Medtronic has notified the Competent Authority of your country of this action. Medtronic remains dedicated to further investigation of this issue and will continue to monitor device performance to ensure we meet your needs and those of your patients. Further communication will follow once more information becomes available. For any additional questions you can reach out to your Medtronic Representative.

Sincerely,

Local / OU Manager