

Did not survive

For Internal Use Only

Clinical Event Reference

Marketing Event Reference

Customer event report

FORM MUST NOT CONTAIN INFORMATION THAT COULD IDENTIFY THE PATIENT

Please do not provide any identifiable information, such as patient name, address or location of hospital.

Patient information						
Male Female Non-binary/ third gender	Age in years:			Weight (estimation):	🗌 Lb	🗌 Kg
Event information						
Country:						
Date of use:		Time of use (local):				
Was the event witnessed?		🗌 Yes	🗌 No	If yes, relationship to patient?		
Was CPR performed by bystander prior to AED switch on?		🗌 Yes	🗌 No	If yes, for how many minutes?		
What was the rescuer response time from SCA to retrieving AED?		In minutes:				
Was patient breathing prior to commencing CPR?		🗌 Yes	🗌 No	Unknown		
Did the patient have a pulse prior to commencing CPR?		🗌 Yes	🗌 No	🗌 Unknown		
Was a shock delivered?		🗌 Yes	🗌 No			
Location type for resuscitation attem	ıpt					

Location type (Check one)	Details
Home	Please indicate the specific type of location (gym, dentist office, restaurant, etc.), providing as much information as possible.
□ Office	DO NOT PROVIDE PLACE NAME, ADDRESS OR GEOGRAPHICAL LOCATION.
Medical facility	
Sports center	
Public space	
Other (Describe location, without name or geographical location)	
Patient outcome	
Outcome (Check one)	Details
Survived to hospital admission	Please provide any additional information on rescue attempt (when did ambulance arrive, actions taken).
Survived to hospital discharge	DO NOT PROVIDE CITY, OR HOSPITAL NAME OR ADDRESS.

Patient pre-existing medical condition (if known)

Condition (Check all that apply)	Please list other known conditions:
🗌 Diabetes mellitus	
Hypertension	
🗌 Hyperlipidaemia	
Implanted pacemaker	

Event file

The event file, downloaded using SAVER EVO software, must be provided with this form. Please use the following filename structure: **Device serial number_Date of event (MM-DD-YYYY)**

Please send both the form and the event file (.evo) to AEDEvent@Stryker.com. A PDF file will not be accepted.

If you need assistance downloading the file, please contact support at HeartSineSupport@stryker.com.

Device information		Pad-Pak [™] information					
Device type (Check one)		Device serial number	Pad-Pak type (Check one) Lot/S		erial number	Expiration date	
SAM PAD 300 SAM	M PAD 360P		🗌 Pad-Pak				
SAM PAD 300P	M PAD 450P		☐ Pediatric-Pak [™]				
SAM PAD 350P	M PAD 500P						
Reporter information		User information					
Event reporter name:			Was user trained? (if known):		Го		
Telephone:			Training provider (if known):			
Email:							
Distributor name:							

Terms

Following are the terms for the Free Pad-Pak and Forward Hearts programs.

- $\label{eq:last} \textbf{l.} \ \ \textbf{Please do not attach any picture, audio and/or video recording related to the reported event.}$
- 2. Event must be a sudden cardiac arrest to qualify. (Event is reviewed by Stryker Clinical team whose decision is final.)
- 3. Please refer to heartsine.com for the complete list of requirements to qualify for Free Pad-Pak and/or Forward Hearts after a Stryker AED has been used during a sudden cardiac arrest resuscitation.

The person completing this form will ensure compliance with local privacy regulations, and agrees to ensure no identifiable information is contained in this form.

Please detail your experience using this AED.

Please do not provide any identifiable information on individuals and places involved.