

HeartSine® samaritan® PAD 350P/360P

Connected AEDs

Semi-Automatic/Fully Automatic Public Access Defibrillators with Integrated Wi-Fi[®] Connectivity

Data sheet

A Ready AED. So simple.

Sudden cardiac arrest strikes with little time to react and even less time to think. This means an Automated External Defibrillator (AED) must be close at hand, easy to use and ready to shock. Designed for simplicity, HeartSine samaritan PAD 350P and 360P Connected AEDs offer key features that help ensure readiness:

- Simplified readiness monitoring
- Integrated Wi-Fi[®] connectivity
- AED program management
- Low cost of ownership

Readiness made easy

- LIFELINKcentral AED Program Manager Monitors AED programs by tracking AED readiness status, Pad-Pak[™] expirations, CPR/AED training certificates and more.
- Integrated connectivity Communicates via Wi-Fi with LIFELINKcentral[™] AED Program Manager to enable AEDs to be managed across a single or multiple locations.



Made for you

Real-time CPR coaching

Easy to understand visual and voice prompts guide the rescuer through the entire resuscitation process, including CPR—a key link in the chain of survival.

- One- or two-button operation With just an ON/OFF button (and the SHOCK button on the SAM 350P), offers a simple, straightforward operation.
- Automatic shock delivery / Motion detection After analysing heart rhythm, automatically delivers a shock (if needed), eliminating the need for the rescuer to push a shock button (SAM 360P).
- Portable and lightweight Most portable AED with its light weight (2.83 lb) and compact footprint.

Clinically validated technology

Proprietary electrode technology and SCOPE[™] biphasic technology, a low energy escalating waveform, that automatically adjusts for differences in patient impedance.

 Highest level of protection against dust and water

Offers unmatched ruggedness with its high IP56 rating.

Simple to own

• Two parts, one expiration date

The innovative Pad-Pak, an integrated battery and electrode single-use cartridge with one expiration date, offers one simple maintenance change every four years.

Low cost of ownership

Shelf life of four years means that the Pad-Pak offers significant savings over other defibrillators that require separate battery and electrode replacements.

8-year Warranty

AED is backed by an 8-year warranty.



Specifications

Defibrillator

Waveform: Self-Compensating Output Pulse Envelope (SCOPE) optimised biphasic escalating waveform compensates energy, slope and duration for patient impedance

Patient analysis system

Method: Evaluates patient's ECG, signal quality, electrode contact integrity and patient impedance to determine if defibrillation is required

Sensitivity/Specificity: Meets IEC/EN 60601-2-4

Impedance range: 20-230 ohms

Energy selection

Pad-Pak shock: Shock 1: 150J Shock 2: 150J Shock 3: 200J

Pediatric-Pak[™]:

Shock 1: 50J Shock 2: 50J Shock 3: 50J

Charge time (typical): 150J in < 8 seconds, 200J in < 12 seconds

Environmental

Operating/Standby temperature: 32°F to 122°F (0°C to 50°C)

Transportation temperature:

14°F to 122°F (-10°C to 50°C) for up to two days. If the device has been stored below 32°F (0°C), it should be returned to an ambient temperature of between 32°F to 122°F (0°C to 50°C) for at least 24 hours before use.

Relative humidity: 5% to 95% non-condensing

Water resistance: IEC 60529/ EN60529 IPX6 with electrodes connected and battery installed

Dust resistance: IEC 60529/ EN60529 IP5X with electrodes connected and battery installed

Enclosure: IEC/EN 60529 IP56

Altitude: 0 to 15,000 feet (0 to 4,575 meters)

Shock: MIL STD 810F Method 516.5, Procedure 1 (40 G's)

Procedure 1

Category 4 Truck Transportation - US Highways

Aviation

EMC: IEC/EN 60601-1-2

Radiated emissions: IEC/EN 55011

Electrostatic discharge: IEC/EN 61000-4-2 (8 kV)

> **RF immunity:** IEC/EN 61000-4-3 80MHz-2.5 GHz. (10 V/m)

Magnetic field immunity: IEC/EN 61000-4-8 (3 A/m)

(Category M)

RTCA/DO-227 (TSO/ETSO-C142a) Falling height: 3.3 feet (1 meter)

Physical characteristics

Size: 9.21 in x 7.25 in x 1.9 in (23.4 cm x 18.4 cm x 4.8 cm)

Weight: 2.83 lb (1.285 kg)

Accessories

(Adult)

Shelf life/Standby life: See the expiration date on the Pad-Pak/Pediatric-Pak (4 years from manufacture date)

Weight: 0.44 lb (0.2 kg) Size:

3.93 in x 5.24 in x 0.94 in (10 cm x 13.3 cm x 2.4 cm)

Battery type: Disposable single-use combined battery and defibrillation electrode cartridge (lithium manganese dioxide (LiMnO2) 18V)

Battery capacity (new): > 60 shocks at 200J or 6 hours of continuous monitoring

are supplied as standard with each device Electrode placement: Anterior - lateral

Anterior - posterior or Anterior - lateral (Pediatric)

Vibration: MIL STD 810F Method 514.5,

Category 7 Aircraft – Jet 737 & General

Aircraft: RTCA/DO-160G, Section 21

With Pad-Pak inserted and HeartSine Gateway[™], with batteries, attached:

Pad-Pak Electrode and Battery Cartridge

Electrodes: Disposable defibrillation pads

Electrode active area: 15 in² (100 cm^2)

Electrode cable length: 3.3 feet (1 meter)

Aircraft safety test (TSO/ETSO-certified Pad-Pak): RTCA/DO-227 (TSO/ETSO-C142a)

HeartSine Gateway Battery

Type: CR123A 3V, Non-rechargeable

Type number: 6205

Designation IEC: CR 17345

Weight (per battery): 17g

Quantity: Four

System: Lithium Manganese Dioxide / Organic Electrolyte

UL recognition: MH 13654 (N)

Nominal voltage (per battery): 3V

Typical capacity load: 100 Ohm, at 68°F (20°C), 1550 mAh down to 2V

Volume: 0.43 in³ (7 ccm)

Data storage

Memory type: Internal memory

Memory storage: 90 minutes of ECG (full disclosure) and event/incident recording

Review: Custom USB data cable (optional) directly connected to PC with Saver EVO™ Windows[®]-based data review software

Materials used

Defibrillator housing / HeartSine Gateway: ABS, Santoprene

Electrodes: Hydrogel, Silver, Aluminium and Polyester

Warranty

AED: 8-year limited warranty

HeartSine Gateway: 2-year limited warranty

Communications

Wireless 802.11 b/g/n data transfer to LIFELINKcentral AED Program Manager or LIFENET System. USB connection to Saver EVO software through Micro USB port

BRIEF SUMMARY OF INDICATIONS AND IMPORTANT SAFETY INFORMATION

INDICATIONS FOR USE

The HeartSine samaritan PAD SAM 350P (SAM 350P), HeartSine samaritan PAD SAM 360P (SAM 360P) and HeartSine samaritan PAD SAM 450P (SAM 450P) are indicated for use on victims of cardiac arrest who are exhibiting the following signs: unconscious, not breathing, without circulation (without a pulse). The devices are intended for use by personnel who have been trained in their operation. Users should have received training in basic life support/ AED, advanced life support or a physician-authorized emergency medical response training program. The devices are indicated for use on patients greater than 8 years old or over 55 lbs (25 kg) when used with the adult Pad-Pak™ (Pad-Pak-01 or Pad-Pak-07). They are indicated for use on children between 1 and 8 years of age or up to 55 lbs (25 kg) when used with the Pediatric-Pak (Pad-Pak-02).

CONTRAINDICATION

If the patient is responsive or conscious, do not use the HeartSine samaritan PAD to provide treatment

WARNINGS

AEDs:

- · The HeartSine samaritan PAD delivers therapeutic electrical shocks that can cause serious harm to either users or bystanders. Take care to ensure that no one touches the patient when a shock is to be delivered.
- · Touching the patient during the analysis phase of treatment can cause interference with the diagnostic process. Avoid contact with the patient while the HeartSine samaritan PAD is analyzing the patient. The device will instruct you when it is safe to touch the patient.
- Do not delay treatment trying to find out the patient's exact age and weight. If a Pediatric-Pak or an alternative suitable defibrillator is not available, you may use an adult Pad-Pak.
- The SAM 360P is a fully automatic defibrillator. When required, it will deliver a shock to the patient WITHOUT user intervention.
- The SAM 450P CPR Rate Advisor is currently only intended to provide feedback on adult patients. If you treat a pediatric patient with the SAM 450P and an adult Pad-Pak, ignore any voice prompts regarding the rate of CPR.
- Do NOT use the HeartSine samaritan PAD in the vicinity of explosive gases, including flammable anesthetics or concentrated oxygen.
- Do NOT open or repair the device under any circumstances as there could be danger of electric shock. If damage is suspected, immediately replace the HeartSine samaritan PAD.

Pad-Paks:

- Do not use if the gel is dry.
- The Pediatric Pad-Pak is not for use on patients under 1 year old. For use with children up to the age of 8 years or up to 55 lbs (25 kg). DO NOT DELAY THERAPY IF YOU ARE NOT SURE OF EXACT AGE OR WEIGHT.
- Only HeartSine samaritan PADs with the 👫 label are suitable for use with the Pediatric-Pak. If the HeartSine samaritan PAD you are using does not have this label, use the adult Pad-Pak if no alternatives are available.
- The use of the Pediatric-Pak will enable delivery of 50J shocks to the pediatric patient.
- The Pediatric-Pak contains a magnetic component (surface strength 6500 gauss). Avoid storage next to magnetically sensitive storage media. It is advised that Pediatric-Paks are stored separately when not in use.
- Never charge, short circuit, puncture, deform, incinerate, heat above 85°C or expose contents of TSO (Aviation) Pad-Pak to water. Remove when discharged.

PRECAUTIONS

AEDs:

- Proper placement of the HeartSine samaritan PAD electrode pads is critical. Electrode pads must be at least 1 in (2.5 cm) apart and should never touch one another.
- Do not use electrode pads if pouch is not sealed.
- Check the device periodically in accordance with the service and maintenance instructions provided in the User Manual.
- Operate the HeartSine samaritan PAD at least 6 feet (2 meters) away from all radio frequency devices or switch off any equipment causing interference.
- · Use of the device outside the operating and storage ranges specified in the User Manual may cause the device to malfunction or reduce the shelf life of the Pad-Pak.
- Do not immerse any part of the HeartSine samaritan PAD in water or any type of fluid.
- Do not turn on the device unnecessarily as this may reduce the standby life of the device.
- Do not use any unauthorized accessories with the device as the HeartSine samaritan PAD may malfunction if non-approved accessories are used.
- Dispose of the device in accordance with national or local regulations.
- Check with the relevant local government health department for information about any requirements associated with ownership and use of a defibrillator in the region where it is to be used.

Pad-Paks:

· Check expiration date.

Saver EVO" Software:

Download the complete HeartSine samaritan PAD memory prior to erasing it. This information should be stored safely for future reference. Ensure that only the events you want to delete have been selected prior to deleting. Once deleted from your computer's memory, events cannot be regenerated and all information will be lost.

POTENTIAL ADVERSE EFFECTS

The potential adverse effects (e.g., complications) associated with the use of an automated external defibrillator include, but are not limited to, the following:

- Failure to identify shockable arrhythmia.
- Failure to deliver a defibrillation shock in the presence of VF or pulseless VT, which may result in death or permanent injury.
- Inappropriate energy which could cause failed defibrillation or post-shock dysfunction.
- Myocardial damage.
- Fire hazard in the presence of high oxygen concentration or flammable anesthetic agents.
- Incorrectly shocking a pulse-sustaining rhythm and inducing VF or cardiac arrest.
- Bystander shock from patient contact during defibrillation shock.
- Interaction with pacemakers.
- Skin burns around the electrode placement area
- Allergic dermatitis due to sensitivity to materials used in electrode construction.
- Minor skin rash.

CAUTION

U.S. Federal law restricts this device to sale by or on the order of a physician.

Please consult the User Manual at www.heartsine.com for the complete list of indications, contraindications, warnings, precautions, potential adverse events, safety and effectiveness data, instructions for use and other important information.

For further information contact us at heartsinesupport@stryker.com or visit our website at www.heartsine.com.

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The HeartSine products CE

described in this brochure meet the European Medical Directive requirement.

UL Classified. See complete marking on product.

H009-043-010-0 US

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