1. **What is the purpose of this advisory circular (AC)?**

This AC provides guidance about onboard emergency medical equipment, including Automated External Defibrillators (AED) and Emergency Medical Kits (EMK). It is intended to guide air carriers when establishing protocols for emergency medical equipment. The Federal Aviation Administration (FAA) expects and anticipates some variation among the programs that air carriers establish for emergency medical equipment. (Also see AC 121-34B, Emergency Medical Equipment Training.)

2. **Does this AC supersede any existing ACs?**

This AC supersedes AC 121-33A, Emergency Medical Equipment, dated May 9, 2003. It also relates to existing AC 120-44A, Air Carrier First Aid Programs (http://www.faa.gov/avr/afs/cabinsafety/acidx.cfm), which is also a good reference source.

3. **What FAA regulations does this AC cover?**


4. **Who should read this AC?**

FAA aviation safety inspectors (cabin safety and operations), part 121 air carrier certificate holders, directors of operations, directors of safety, crewmembers, AED manufacturers and suppliers, EMK suppliers, as well as people involved in the development of air carrier procedures and training programs. This AC may also be valuable to people associated with operations under 14 CFR part 125, part 135, and subpart K of part 91 (fractional ownership programs).

5. **When is an emergency medical kit and an AED required and on what size of aircraft?**

The FAA requires AEDs on all airplanes of air carriers operating under part 121 with a maximum payload capacity of more than 7,500 pounds and with at least one flight attendant. Affected airplanes typically would have a capacity for 30 passengers or more requiring at least one flight attendant. The FAA also requires an EMK on all airplanes of air carriers operating...
under part 121 for which at least one flight attendant is required. EMKs and AEDs are “no-go” items and must be carried as indicated on the Minimum Equipment List.

6. **What emergency medical equipment must air carriers carry?**

At least one approved AED, legally marketed in the United States in accordance with Food and Drug Administration (FDA) requirements.

At least one approved EMK with the following items.

Part 121, appendix A, specifies that the following items must be carried in EMKs:

<table>
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<th>CONTENTS</th>
<th>QUANTITY</th>
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<td>Sphygmomanometer</td>
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<tr>
<td>Stethoscope</td>
<td>1</td>
</tr>
<tr>
<td>Airways, oropharyngeal (3 sizes): 1 pediatric, 1 small adult, 1 large adult or equivalent</td>
<td>3</td>
</tr>
<tr>
<td>Self-inflating manual resuscitation device with 3 masks (1 pediatric, 1 small adult, 1 large adult or equivalent)</td>
<td>1: 3 masks</td>
</tr>
<tr>
<td>CPR mask (3 sizes), 1 pediatric, 1 small adult, 1 large adult, or equivalent</td>
<td>3</td>
</tr>
<tr>
<td>IV Admin Set: Tubing w/ 2 Y connectors</td>
<td>1</td>
</tr>
<tr>
<td>Alcohol sponges</td>
<td>2</td>
</tr>
<tr>
<td>Adhesive tape, 1-inch standard roll adhesive</td>
<td>1</td>
</tr>
<tr>
<td>Tape scissors</td>
<td>1 pair</td>
</tr>
<tr>
<td>Tourniquet</td>
<td>1</td>
</tr>
<tr>
<td>Saline solution, 500 cc</td>
<td>1</td>
</tr>
<tr>
<td>Protective nonpermeable gloves or equivalent1</td>
<td>1 pair</td>
</tr>
<tr>
<td>Needles (2-18 ga., 2-20 ga., 2-22 ga., or sizes necessary to administer required medications)</td>
<td>6</td>
</tr>
<tr>
<td>Syringes (1-5 cc, 2-10 cc, or sizes necessary to administer required medications)</td>
<td>4</td>
</tr>
<tr>
<td>Analgesic, non-narcotic, tablets, 325 mg</td>
<td>4</td>
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<tr>
<td>Antihistamine tablets, 25 mg</td>
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</tr>
<tr>
<td>Antihistamine injectable, 50 mg, (single dose ampule or equivalent)</td>
<td>2</td>
</tr>
<tr>
<td>Atropine, 0.5 mg, 5 cc (single dose ampule or equivalent)</td>
<td>2</td>
</tr>
<tr>
<td>Aspirin tablets, 325 mg</td>
<td>4</td>
</tr>
<tr>
<td>Bronchodilator, inhaled (metered dose inhaler or equivalent)</td>
<td>1</td>
</tr>
<tr>
<td>Dextrose, 50%/50 cc injectable, (single dose ampule or equivalent)</td>
<td>1</td>
</tr>
<tr>
<td>Epinephrine 1:1000, 1 cc, injectable, (single dose ampule or equivalent)</td>
<td>2</td>
</tr>
<tr>
<td>Epinephrine 1:10,000, 2 cc, injectable, (single dose ampule or equivalent)</td>
<td>2</td>
</tr>
<tr>
<td>Lidocaine, 5 cc, 20 mg/ml, injectable (single dose ampule or equivalent)</td>
<td>2</td>
</tr>
<tr>
<td>Nitroglycerine tablets, 0.4 mg</td>
<td>10</td>
</tr>
<tr>
<td>Basic instructions for use of the drugs in the kit</td>
<td>1</td>
</tr>
</tbody>
</table>

1 Although the FAA requires only one pair of protective gloves, it recommends that operators keep additional pairs accessible on the aircraft. This would allow crewmembers to access a pair of gloves without having to locate and open an EMK.
7. What is the purpose of the following items contained in the EMK?

- **Non-narcotic analgesic tablets**: a general oral medication used mainly to relieve muscle aches and headaches
- **Oral antihistamine**: medication used mainly to relieve symptoms associated with allergies and hay fever
- **Aspirin**: a general oral medication used mainly to alleviate head and muscle aches and chest pain or heart attack
- **Atropine**: medication used mainly to increase heart rate, that may be needed to assist a passenger with an unstable cardiac rhythm
- **Bronchodilator inhaler**: a preparation of medication used to help restore normal breathing in asthmatics
- **Epinephrine 1:10,000**: medication used mainly for cardiac resuscitation
- **Lidocaine**: medication used mainly in cases of unresponsiveness to defibrillation and possibly for maintenance of normal heart rhythm after successful defibrillation
- **An IV administration set including tubing with 2Y connectors (and, for placing the IV, alcohol sponges, tape, bandage scissors, and a tourniquet)**: equipment used for administering IV drugs (e.g., atropine, lidocaine, epinephrine) that may be needed to sustain heart function
- **A self-inflating manual resuscitation bag (AMBU bag) (with 3 masks: 1 pediatric, 1 small adult, and 1 large adult)**: equipment that may be needed for continuation of respiratory support
- **CPR mask (1 pediatric, 1 small adult, 1 large adult)**: equipment that may be needed to protect a person while administering CPR

8. What does “or equivalent” mean?

The FAA recommends that air carriers carry the required EMK items without substitution. The FAA has used the words “or equivalent” in part 121, appendix A, since 1986 (and will continue to use the words) to allow for any nomenclature or other changes the medical community might choose to make over the course of the lifetime of the regulation. The FAA references only generic terms under part 121, appendix A as amended. If you have a question about whether a certain medication or piece of equipment you choose to stock will meet the requirement, please contact the FAA Office of Aerospace Medicine.

Suppliers have asked the FAA whether diphenhydramine HCl injection is an acceptable equivalent to meet the requirement for antihistamine injectable. It is acceptable. They also have asked whether it is acceptable to stock universal masks where CPR masks or masks for resuscitation are required. In both situations, universal masks designed for the required sizes are acceptable as long as they meet the quantity requirements. In addition, some masks may be used to administer CPR and also may be used with the self-inflating manual resuscitation device. These masks often use a one-way valve, to protect the rescuer during CPR, and a separate connector for the resuscitation device. If the universal masks included in the EMK provide a
means of administering CPR and also may be used with the self-inflating manual resuscitation device, then they are considered to be acceptable under both mask requirements. Therefore, a total of only three masks would be required.

9. What does “approved” EMK and “approved” AED mean?

Approved EMK means that the FAA Principal Operations Inspector assigned to the holder of an operating certificate exercises approval for the Administrator, as appropriate, of equipment to be carried aboard a certificate holder’s aircraft.

Approved AED means that it is legally marketed in the United States in accordance with FDA requirements. AEDs used on airplanes must be approved by the FDA for medical use and must conform to FDA standards.

10. How can an air carrier comply with part 121, appendix A, at all times after an EMK and/or an AED is used during flight?

The regulation specifies “at least one” EMK and “at least one” AED as the minimum required on every flight for full compliance with part 121, appendix A. In the event that certain contents of an EMK are used during a flight, an inventory of the remaining contents and restocking of the contents would be needed to ensure that the minimum content requirements are met prior to any subsequent flight. For the sake of convenience, and to avoid delays, an airline may decide to overstock certain EMK items (in particular protective gloves and CPR masks), carry two EMKs, or establish a procedure for effecting one-for-one replacements as necessary.

An air carrier may elect to carry redundant equipment to ensure that after use of equipment in flight, the minimum required equipment is still on board for dispatch. In such circumstances flight attendants need to be aware of any inoperative AEDs or incomplete EMKs in the cabin in order to avoid the possibility that during an inflight medical emergency someone tries to use an inoperative AED or searches for a missing item in an incomplete EMK. In order to make flight attendants aware of inoperative equipment, an air carrier may consider the following effective practices:

- Labeling inoperative AEDs with a statement such as “Inoperative – Do Not Use”
- Labeling incomplete EMKs with a statement such as “Incomplete – Missing Contents”
- Implementing a procedure (briefing) that ensures all flight attendants are aware of incomplete EMKs or inoperative AEDs in the aircraft cabin

But, as previously noted in paragraph 5, if the air carrier elects to have only one AED and one EMK on board, if that AED is inoperative or that EMK is incomplete, the aircraft may not be dispatched.

The FAA also acknowledges that there may be circumstances that would warrant a flight attendant needing only protective gloves, a CPR mask, or both from the EMK. Accessing an
EMK for the purpose of retrieving one or both of these items could be problematic. Therefore, the FAA recommends that air carriers carry a few pairs of extra protective gloves and an extra CPR mask outside of the EMK.

The issue of AED replacement will not be as critical as EMK replacement unless, for example, an air carrier allows an AED to be taken off their aircraft for continued assistance of a passenger during emergency ground transport. Individual airlines should develop a protocol for AED use, post-resuscitation guidelines, and any AED serviceability needs. At a minimum, before any subsequent flights, the AED must be “operative” and there must be at least one set of unused pads with the AED. AEDs usually are packaged with a spare battery and a spare set of pads. Air carriers may want to carry extra AED pads.

11. Who is allowed to use the equipment?

Flight attendants should grant access to the equipment only to trained crewmembers or to other persons qualified and trained in the use of emergency medical equipment. The decision to allow passengers to assist another passenger and have access to medical equipment is up to the air carrier and its agents. The FAA does not attempt to define the various medical specialties under part 121 because it limits access to the extent that the only person available to assist on a flight might not be included. It would be preferable for flight attendants to check the credentials of passengers holding themselves out as medical specialists.

It is unrealistic to expect flight attendants to achieve the same level of proficiency as emergency medical personnel who perform medical procedures on a routine basis. Flight attendants should not be expected to administer medications or to start IVs. If a critical in-flight medical event occurs and a passenger medical specialist is not available, it is recommended that the sick passenger be made as comfortable as possible and the pilot in command should determine whether to attempt safe diversion of the aircraft.

As stated in the rule, the decision to offer treatment or take other action (including safe diversion of the aircraft) is discretionary with the air carrier and its agents. The FAA does not require any actions by the air carrier and its agents and/or other passengers other than having certain emergency medical equipment on board the aircraft.

12. What does “readily accessible” mean under § 121.803?

In § 121.803, the FAA uses the term “readily accessible” in the same way as the longstanding terminology used for all emergency equipment under § 121.309 (b)(2). “Readily accessible” means, as it always has, that air carriers should place equipment where crewmembers can access the equipment quickly. “Readily accessible” is not intended to mean that the emergency medical equipment should be located where it might be subject to unauthorized access.
13. Where should we store this equipment?

Because of the various configurations of aircraft, the FAA does not set one standard for storing the equipment. Airlines typically put the equipment in a locked compartment in an overhead bin, in a locked compartment attached to the bulkhead behind the last row of seats or in first class, or in an unlocked pouch attached to a bulkhead behind the last row of seats. All of these methods are acceptable. To avoid unnecessary distraction on the flight deck, and to ensure flight deck integrity, do not store AEDs in flight deck compartments.

14. How must we inspect the equipment?

You must regularly inspect emergency medical equipment in accordance with inspection periods established in your operations specifications and maintain it according to manufacturers’ specification. You should follow the manufacturer’s recommended procedures regarding an AED self-check.

Flight attendants perform a routine preflight inspection of all emergency medical equipment in accordance with their air carrier’s procedure to assure that it is on board the aircraft, secured, and ready if needed for use. Since EMKs are sealed, it’s difficult to do a comprehensive visual inspection to ascertain that no EMK items are missing or unusable; therefore, it is critical to assure EMK integrity prior to the preflight inspection stage. Any discrepancies must be resolved in accordance with your air carrier’s procedures.

15. Most self-inflating manual resuscitation devices (AMBU bags) found in an EMK are accompanied by tubing that can be connected to an outlet on a portable oxygen bottle located in the aircraft cabin. This allows additional pure oxygen to mix with the ambient air in the AMBU bag and raises the level of oxygen provided during a medical event where the AMBU bag is used for respiratory support. Is this practice permissible?

Yes. Current regulations do not prohibit the connection or disconnection of oxygen masks and/or tubing that is provided with the AMBU bag in the EMK to an outlet on the regulator of an air carrier’s portable oxygen bottle during a medical event that occurs in flight.

16. How often should we replace the EMK items?

The medications that must be carried in all EMKs have an expiration date of approximately 1 year: atropine, bronchodilator inhaler, dextrose, epinephrine, saline solution, and lidocaine; aspirin, non-narcotic analgesic, antihistamine, and nitroglycerine tablets. If temperature extremes occur on the aircraft at any time or if the medications have surpassed their expiration date then you should replace them. The FAA has not found expiration of medications to be problematic for air carriers under the existing requirement to carry injectable antihistamine, dextrose, epinephrine, and nitroglycerin tablets. Therefore, the FAA does not anticipate that replacing medications would become problematic by requiring additional medications of similar shelf-life. The best practice, under normal circumstances, is to replace all of the medications annually.
17. What does “damaging temperatures” mean under part 121, appendix A?

“Damaging temperatures” means temperature extremes which could alter the effectiveness of the emergency medical equipment.

Current manufacturers’ specifications indicate that medications required for the EMK stored at controlled room temperature should remain stable within a temperature range of 59 to 86 degrees Fahrenheit (15 to 30 degrees Celsius). Medications carried in emergency medical vehicles, such as ambulances, reportedly remain stable within an even wider temperature range. The EMK and the aircraft cabin provide some protection from potentially harmful external temperatures. The aircraft cabin environment does not appear to negatively affect the required medications as long as they are replaced before their expiration date.

If an aircraft has been exposed to extremes of hot or cold temperatures, the medications in a liquid form (injectable) should be inspected before use. If they are yellow or cloudy, then they may have lost their effectiveness and should not be used. In general, once injectable medications are frozen they should not be used, and high, prolonged heat will degrade the efficacy of most medications.

In addition, the AED, batteries, and defibrillator pads usually have a recommended temperature range for storage and operation. These temperature ranges vary between manufacturers, but are generally much wider than for the medications. The manufacturers’ specifications should be consulted for proper handling procedures if the aircraft cabin exceeds the recommended temperatures. Prolonged exposure to temperatures outside the recommended limits may damage the batteries or may cause the pads to not adhere properly.

If an aircraft is parked or taken out of service for an extended period of time in a location where it may be exposed to temperature extremes, then the emergency medical equipment should be taken off the aircraft and protected.

18. Since some air carriers carry EMKs that may contain controlled substances, how can they be transported legally? Is transporting these substances compatible with Drug Enforcement Administration (DEA) regulations?

Although the FAA does not require any controlled substances for the EMK, some air carriers may purchase commercial EMKs that are prepackaged with a controlled substance(s) (for example, diazepam). Such EMKs cannot be purchased (or carried) unless a current DEA Registration Certificate is on file with the EMK distributor. If a controlled substance is compromised (e.g., lost, stolen, or missing) the air carrier must report it to the DEA.

19. Does the FAA regulate safety standards for AEDs?

No. The FDA is responsible for regulating safety standards for the manufacture and use of AEDs. The FAA is responsible for regulating the safety of the power sources used in AEDs when carried on board a passenger-carrying aircraft. You should direct any questions about
AEDs directly to the manufacturer and/or to the FDA Center for Devices and Radiological Health. AED manufacturers may have resources available to provide the FDA-required oversight.

For safety purposes, the FAA asks that certificate holders comply with the guidance in applicable Flight Standards Information Bulletins for Airworthiness, such as FSAW 98-05, Medical Portable Electronic Devices (PED). Certificate holders must also comply with the requirements of applicable FAA Technical Standard Orders (TSO) such as TSO-C142, Lithium Batteries. The devices should be maintained in accordance with manufacturers’ specifications and should be inspected in accordance with schedules developed under operations specifications. Currently, AEDs are powered by primary (not rechargeable) lithium batteries. Safety of these batteries is stressed because extremely energetic materials are used in lithium cells and they are not intrinsically safe. Safety concerns include the possibility of fire, explosion, and the venting of toxic or flammable gases.

20. What are acceptable power sources for AEDs?

The FAA requires the power source (e.g., batteries) used to power AEDs to comply with all requirements in applicable advisory material such as Advisory Circular 91.21-1A Use of Portable Electronic Devices Aboard Aircraft (http://www.airweb.faa.gov/Regulatory_and_Guidance_Library/rgAdvisoryCircular.nsf/MainFrame?OpenFrameSet ), and in applicable TSOs, such as TSO-C142, Lithium Batteries (http://avinfo.faa.gov/tso/tsocur/current.htm).

On March 24, 2005, the FAA amended the regulations for emergency medical equipment to allow approved power sources that do not have TSO markings to be used in AEDs carried onboard aircraft (http://dmses.dot.gov/docimages/p80/322276.pdf). AED power sources manufactured before July 30, 2004, and not TSO marked, may continue to be used until their expiration date, provided that the power source manufacturer has requested and received from the FAA a finding of TSO equivalency for its product.

Specifically, part 121, Appendix A, was amended to allow the use of AED power sources that were manufactured before July 30, 2004, and do not have the TSO marking required, provided that the manufacturer of the power source has received a finding of equivalency from the appropriate Aircraft Certification Office (ACO). A manufacturer can seek this determination by contacting the ACO that issued the TSO approval of its AED power source.

21. Is labeling an AED with the statement “approved for use on aircraft” appropriate?

No. Arbitary use of the statement "approved for use on aircraft" is not appropriate. It could lead to a safety problem because toxic gas venting precautions are required before placing AEDs containing lithium sulfur dioxide batteries in an airplane cabin. The battery manufacturer must supply a note with the batteries that addresses installation procedures and limitations. Marking requirements for lithium batteries are defined in part 21, specifically § 21.607(d).
22. What prompted requirements for emergency medical equipment?

The Aviation Medical Assistance Act (the Act) of 1998 [Pub. L. 105-170, 49 U.S.C. 44701] directed the FAA to determine whether current minimum requirements for air carrier emergency medical equipment and air carrier crewmember emergency medical training should be modified. As directed in the Act, the FAA conducted a year-long data collection on death or near-death in-flight medical events. It revealed 188 total events resulting in 108 deaths (119 of these 188 total incidents were cardiac-related resulting in 64 deaths). For cardiac-related events on the aircraft, an AED was reported as “not available” for 40 events. An AED was available and used to deliver at least one shock in 17 separate events. From these events, four passengers were reported as having survived. Subsequent to the data collection, further investigation revealed that more passengers, and a flight crewmember, had also survived after having been shocked with an AED. Based on these events, it was determined that part 121 should be amended to require emergency medical enhancements, such as performance-based training for flight attendants on the use of AEDs and CPR, enhanced EMKs, and AEDs.

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