

WHO/PQS/E03/RF05.1 Original: English Distribution: General

TITLE: Refrigerator or combined refrigerator-icepack freezer: compression-cycle. Solar direct drive without battery storage

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1. Scope:

This specification defines the requirements for compression-cycle vaccine refrigerators or combined vaccine refrigerator/icepack freezers powered by a solar electric system with no battery. Pre-qualification testing will establish a minimum solar radiation reference period below which the product should not be used and the maximum autonomy that the product can achieve. PQS specification **E03/PV01** specifies a compatible **Type 2** solar power system. Three temperature zone designations are described: moderate zone, temperate zone and hot zone. In addition appliances are tested to establish a minimum rated ambient temperature designation.

2. Normative references:

DIN 8985: 1983-05: Testing the surfaces of installed refrigerators and freezers.

EMAS: European Union Eco-Management and Audit Scheme.

IEC 60335-1: Amendment 1 - Household and similar electrical appliances - Safety - Part 1: General requirements.

IEC 60335-2-24: 2007 -Household and similar electrical appliances -Safety -

Part 2-24: Particular requirements for refrigerating appliances, ice-cream appliances and ice-makers.

ISO 9001: 2000: Quality Management Systems – Requirements.

ISO 14001: 2004: Environmental management systems - Requirements with guidance for use.

ISO/IEC 17025: 2000: General requirements for the competence of testing and calibration laboratories.

ISO 20282-1: 2006: Ease of operation of everyday products - Part 1: Context of use and user characteristics.

WHO/PQS/E03/PV01.1: Solar power system for compression-cycle vaccine refrigerator or combined refrigerator-icepack freezer.

3. Terms and definitions:

Acceptable temperature range: The acceptable temperature range for storing vaccine is $+2^{\circ}$ C to $+8^{\circ}$ C. However, brief excursions outside these limits will be tolerated. Measured over a five day period of testing these excursions must not exceed any of the following parameters:

- The cumulative total of all excursions over $+8^{\circ}$ C must not exceed 12 hours.
- No excursion must exceed +20°C.
- The cumulative total of all excursions below +2°C must not exceed 12 hours.
- No excursion must reach 0°C.

To establish compliance with these requirements, the total hours of runtime for the *stable running*, *icepack freezing*, *day/night cycle* and *minimum rated ambient temperature* tests will be recorded. Temperature logger data will be analyzed to establish the cumulative number of hours with temperatures lying outside the acceptable temperature range. The results will then be normalized to establish the extent of excursions over an average five day period.

Autonomy: Time in days that a solar refrigerator, or combined refrigerator and icepack freezer, can maintain the vaccine load within the acceptable temperature range under low solar radiation conditions (e.g. rain). Autonomy is determined as described in E03/PV1 – Section 4.1.2 and measured as described in E03/RF05-VP.

Holdover time: The time in hours without solar energy input during which all points in the vaccine compartment remain between $+2^{\circ}$ C and $+10^{\circ}$ C without solar energy input and at the maximum ambient temperature of the temperature zone for which the appliance is rated. The same holdover time, or better, must also be achieved at the minimum rated ambient temperature.

Hot zone: Hot zone units must operate at a steady +43°C ambient temperature and over a+43°C/+25°C day/night cycling temperature range.

In writing: means communication by letter, fax or email.

Installation: The refrigerator specified in this document, connected to a solar power system complying with specification **E03/PV01**.

Legal Manufacturer: The natural or legal person with responsibility for the design, manufacture, packaging and labelling of a product or device before it is placed on the market under his own name, regardless of whether these operations are carried out by that person himself or on his behalf by a third party.

Minimum rated ambient temperature: In addition to the day/night test, all appliances will be challenged by reducing the ambient temperature in 5°C increments below the lower limit for the model's rated temperature zone, down to a minimum of -10°C. This test is designed to determine the lowest constant ambient temperature at which the acceptable temperature range can be maintained

with a full vaccine load. Once established, this figure will be displayed in the blue sector of the Annex 1 temperature zone symbol. This will enable purchasers incountries with low winter temperatures to select the most appropriate models. Moderate zone: Moderate zone appliances must operate at a steady +27°C ambient temperature and over a+27°C/+10°C day/night cycling temperature range.

Montreal Protocol: Montreal Protocol on Substances that Deplete the Ozone Layer.

Reseller: A commercial entity, licensed to act on behalf of a Legal Manufacturer, and which carries product liability and warranty responsibilities no less onerous than those carried by the Legal Manufacturer.

Solar Radiation Reference Period: The minimum average daily solar radiation on the plane of the solar array that is required to properly power the solar refrigerator, or combined refrigerator and icepack freezer, expressed in kWh/m²/day.

Temperate zone: Temperate zone units must operate at a steady $+32^{\circ}$ C ambient temperature and over a $+32^{\circ}$ C/ $+15^{\circ}$ C day/night cycling temperature range.

4. Requirements:

4.1 *General*:

Solar powered DC compression vaccine refrigerators are used primarily in areas without any electricity or where there is less than 8 hours of reliable electricity over a typical day. Reliability, durability and effective maintenance is essential for a successful installation. The associated power system must be designed to match both the refrigerator power consumption and local climate conditions (i.e. ambient temperatures and solar radiation resource). Manufacturers may offer products suitable for one or more temperature zones.

4.2 <u>Performance:</u>

4.2.1 *Operating temperature range:*

As indicated on the temperature zone rating sticker attached to the product (see Annex 1).

4.2.2 Refrigeration cycle:

Compression-cycle unit, with one or two compressors, operating on direct current electricity.

4.2.3 Voltage and frequency:

Generally 12 volt DC, supplied from a photovoltaic array.

4.2.4 *Icepack freezing capacity:*

In combined units with freezer compartment, a minimum of 1.6 kg and not less than 2.4 kg per 50 litres of gross freezer volume must be frozen per 24 hours whilst maintaining the temperature control specified in 4.2.5.

4.2.5 Freezing compartment capacity in combined units:

No standard set, but it is recommended that the freezer compartment holds a minimum of 8 kg of icepacks complying with PQS section **E05**. These may be frozen gradually over several days.

4.2.6 Areas not suitable for vaccine storage:

Areas of an otherwise acceptable appliance which are too warm or too cold must be excluded from use by design.

4.2.7 Temperature control:

Refrigerator compartment: The entire vaccine load must remain within the acceptable temperature range during any continuous ambient temperature test(s) or day/night cycling temperature test(s). Combined units must achieve this performance with or without icepacks in the icepack compartment.

Icepack freezing compartment: The icepack freezing compartment (if present) must remain below -5°C under the same ambient conditions.

4.2.8 Thermostat:

The thermostat must be set to prevent freezing in any part of the vaccine storage compartment. The thermostat must be effective throughout the ambient operating temperature range (including the minimum rated ambient temperature range where applicable – see clause 4.2.10). It must be designed so that it cannot be adjusted by the user. A means for adjustment by a technician is acceptable provided the device is protected from user interference (e.g. by location within the appliance cabinet). Alternatively, programmable thermostats may be password-protected.

4.2.9 Thermometer:

- **Option A:** Externally readable cabinet-mounted gas or vapour pressure dial thermometer.
- **Option B:** Externally readable cabinet-mounted electronic maximum-minimum thermometer conforming to PQS specification E06/TH06, powered off the battery set. Dry cell batteries must not be used.

4.2.10 Autonomy:

A minimum of 5 days at the specified solar radiation reference period and temperature. *Note:* this standard will be reduced to 3 days when an autonomy sizing method is provided.

4.2.11 Holdover time:

Holdover time must be a minimum of 20 hours.

4.2.12 Minimum rated ambient temperature:

All models will be tested to establish their minimum rated ambient temperature. The minimum acceptable performance rating is achieved if the product passes the day/night test for its nominal temperature zone. The maximum performance rating is achieved if the vaccine load remains within the acceptable temperature range at -10°C. A freeze-prevention circuit may be required to protect against freezing at low ambient temperatures.

4.2.13 Power consumption:

No standard set; however performance data will be published.

4.2.14 Defrost switch:

If used, the defrost switch (or switches if dual compressors are employed) must be accessible to the user without tools but must be protected from accidental changes in position. No other power switch is to be installed.

4.2.15 Lock:

The door or lid must be fitted with a lock. Two keys are to be supplied with every unit.

4.2.16 Corrosion resistance:

Internal and external cabinet, lid and frame protected against corrosion to DIN 8985.

4.2.17 Electrical safety rating:

Manufacturer to certify compliance with IEC 60335-1 and IEC60335-2-24.

4.2.18 Markings:

Compressors operating on R 134a refrigerant must be marked with the blue identifying symbol shown in Annex 2. In addition, the freezer cabinet must be permanently marked, near the compressor position, with the chemical name of the refrigerant, or with the refrigerant number, formula or proportion (for blended refrigerants).

4.2.19 Vaccine storage advice:

All units must carry a factory-fitted non-removable label, designed to last the lifetime of the appliance, carrying the following information:

- **Vaccine freezers:** Vaccine storage instructions and the appropriate temperature zone symbol as Annex 1.
- **Combined freezers:** Vaccine storage instructions, icepack freezing instructions and the appropriate temperature zone symbol as Annex 1.

The instructions should be fixed to the lid of chest refrigerators and near the top of the door on upright refrigerators. Instructions should be in one of the languages specified in clause 4.11, as indicated by the purchaser at the time of ordering. If removable baskets are supplied fix a multi-lingual warning within the refrigerator instructing users to *Store vaccine in baskets only* or other appropriate instruction.

4.3 *Environmental requirements:*

4.3.1 Ambient temperature range during transport and storage:

 -30° C to $+55^{\circ}$ C when the product is inactivated.

4.3.2 Ambient humidity range during transport, storage and use: 5% to 95% RH.

4.4 *Physical characteristics:*

4.4.1 Overall dimensions:

To allow for manoeuvring through corners, corridors and doorways, the minimum dimension of the product (either length, width or height) should not exceed 830mm, and preferably it should not exceed 710mm. The maximum dimension must not exceed 1700mm and the maximum diagonal (corner to corner) dimension must not exceed 1850mm.

4.4.2 Weight:

Not critical.

4.5 *Interface requirements:*

4.5.1 Battery charge regulator compatibility:

All electrical components must be compatible with a **Type 2** solar power system as specified in specification **E03/PV01**.

4.5.2 Alarm:

The alarm system may be supplied with the photovoltaic power system. However, if it is incorporated into the refrigerator casing, the control panel must have a colour-coded voltmeter, LEDs, or other indicators to show:

- Array charging (green).
- Low battery (orange or yellow), with a clearly labelled warning: *Do not freeze icepacks* in the appropriate language.
- Load disconnect (red).

An acoustic alarm may be included as a supplementary high/low battery and/or load disconnect warning device.

4.6 *Human factors:*

4.6.1 Generally:

The product must be useable by the widest practicable range of active health workers, regardless of age, gender, size or minor disability, including long-sighted people without glasses, in accordance with the general principles laid out in ISO 20282-1: 2006.

4.6.2 *Control panel and thermometer:*

The control panel must be positioned on the front of the unit. The defrost switch, if present, should be recessed or otherwise protected so that it is not possible inadvertently to activate it. The thermometer dial must be positioned at the front of the unit and as close to eye level as possible within the constraints of the cabinet design.

4.7 *Materials*:

4.7.1 Refrigerant:

HFC (hydro fluorocarbon) or HC (hydrocarbon) refrigerant. CFC (chlorofluorocarbon) and HCFC (hydrochlorofluorocarbon) gases are not acceptable. The suitability of alternative refrigerant gases will continue to be assessed.

4.7.2 Thermal insulation foaming agents:

Any gas complying with the limitations and deadlines set by the Montreal Protocol on the elimination of ozone-depleting chemicals.

4.7.3 Other restricted materials:

The product and its constituent components, must not contain lead (except in batteries), mercury, cadmium, hexavalent chromium, polybrominated biphenyls (PBB) or polybrominated biphenyl ethers (PBDE).

4.8 *Warranty:*

The product is to be covered by a two year replacement warranty in the event of any component failure arising from defective design, materials or workmanship.

4.9 <u>Servicing provision:</u>

The product is to be designed to achieve a low-maintenance life of not less than 10 years apart from routine de-frosting, cleaning, solar array shading prevention, battery water addition and replacement of batteries in the associated battery set.

4.10 <u>Disposal and recycling:</u>

The manufacturer is to provide information to the buyer on the hazardous materials contained within the system and suggestions for resource recovery/recycling and/or environmentally safe disposal. For the European Union WEEE compliance in accordance with European Union Directive 2002/96/EC is mandatory.

4.11 *Instructions*:

User and maintenance instructions in Arabic, English, French, Mandarin Chinese, Russian and Spanish. The instructions are to be written for users and repair technicians and are to cover the following topics:

- installation procedures;
- compatible types of solar power system;
- prevention of vaccine freezing;
- simple daily, weekly and monthly maintenance tasks;
- periodic preventative maintenance checks;
- diagnostic and repair procedures;
- itemized list of spare parts including part numbers;
- end-of-life resource recovery and recycling procedures.

4.12 *Training:*

Not required.

4.13 *Verification:*

In accordance with PQS Verification Protocol E03/RF05-VP.

5. Packaging:

Materials used for packaging the finished product are to be free of ozone-depleting compounds as defined in the Montreal Protocol. The general specification of shipping containers will be subject to agreement with the individual procurement agencies.

6. On-site installation:

Not required.

7. Product dossier:

The legal manufacturer or reseller is to provide WHO with a pre-qualification dossier containing the following:

- Dossier examination fee in US dollars.
- General information about the legal manufacturer, including name and address.
- Unique identification reference for the product type.

- Brand name of the product.
- Full specifications of the product being offered, covering all the requirements set out in this document, including details of product marking and traceability.
- Full details of the recommended compatible solar power system (see specification **E03/PV01**).
- A comprehensive set of photographs showing all external surfaces of the unit, the interior layout, the compressor and a close-up of the thermometer and the control panel.
- Certified photocopies of all type-approvals obtained for the product, including CE marking and the like.
- Certified photocopies of the legal manufacturer's ISO 9001: 2000 or more recent quality system certification. Note: Companies that are not ISO 9001-certified will be granted a period of 18 months following the date of publication of this document in which to achieve certification.
- Where relevant, certified photocopies of the legal manufacturer's ISO 14001:2004 certification, EMAS registration or registration with an equivalent environmental audit scheme. Conformity with an environmental audit scheme is not mandatory; however preference will be given to manufacturers who are able to demonstrate compliance with good environmental practice.
- Where available, laboratory test report(s) proving conformity with the product specifications.
- Indicative cost of the product per unit, per 10 units and per 100 units, EXW (Incoterms 2000).

8. On-site maintenance:

Maintenance will be carried out by the end-user and/or his agents.

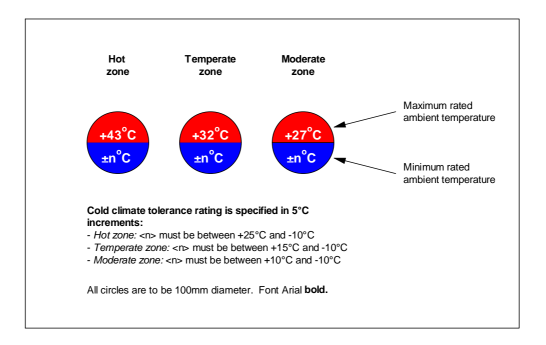
9. Change notification:

The legal manufacturer or reseller is to advise WHO in writing of any changes which adversely affect the performance of the product after PQS pre-qualification has taken place. Any change that WHO considers would alter the test results obtained against the PQS verification protocol **E03/RF05-VP** will result in a request for the product to be retested.

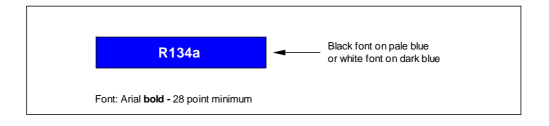
10. Defect reporting:

The legal manufacturer or reseller is to advise WHO and the UN purchasing agencies in writing in the event of safety-related product recalls, component defects and other similar events. If requested to do so by WHO/UNICEF, the manufacturer is to submit a report to WHO/UNICEF stating the number of affected systems and the number of component repairs/replacements provided, together with copies of any associated field reports.

Annex 1 – Temperature zone symbol for refrigerators



Annex 2 - R134a symbol



Revision h	istory:		
Date	Change summary	Reason for change	Approved
05.03.2007	General edit with additional clauses plus Annexes 1 and 2.	Final revisions to PQS format.	UK
09.05.2007	Revised to SMc comments & teleconference UK, SMc, AG 26.04.07		UK
16.05.2007	Final review version		UK
23.05.2007	4.2.6: New clause. 4.2.8: Minor correction 4.11: Minor addition	Consistency with other specs.	UK
08.08.2007	Autonomy definition clarified. 4.2.5: 'No' added before 'standard'. 4.2.10: '3 days' changed to '5 days'. Note re autonomy sizing method added. 4.2.14: 'but must be protected from accidental changes in position' added. 4.2.15: Lock spec changed. 4.4.1: Dims clarified. 4.5.2: 'LEDs or other indicators' added. 4.5.2: Last sentence: 'low' changed to 'high/low'.	Response to manufacturer's comment.	UK