



SLOVENSKI STANDARD
SIST EN 1789:2020+A1:2024

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Medicinska vozila in pripadajoča oprema - Cestna reševalna vozila

Medical vehicles and their equipment - Road ambulances

Rettungsdienstfahrzeuge und deren Ausrüstung - Krankenkraftwagen

Véhicules de transport sanitaire et leurs équipements - Ambulances routières

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43.160 Vozila za posebne namene Special purpose vehicles

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Medical vehicles and their equipment - Road ambulances

Véhicules de transport sanitaire et leurs équipements -
Ambulances routièresRettungsdienstfahrzeuge und deren Ausrüstung -
Krankenkraftwagen

This European Standard was approved by CEN on 13 April 2020 and includes Amendment approved by CEN on 21 September 2023.

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Contents

Page

European foreword.....	4
Introduction	5
1 Scope	6
2 Normative references	6
3 Terms and definitions	9
4 Requirements	10
4.1 General requirements	10
4.2 Electrical requirements	10
4.2.1 General.....	10
4.2.2 Electromagnetic compatibility (EMC).....	11
4.2.3 Battery and alternator	11
4.2.4 Electrical installation.....	12
4.2.5 Visual warning system and audible warning system (siren)	13
4.2.6 Reversing systems	13
4.2.7 Exterior illumination lights.....	13
4.3 Vehicle body	14
4.3.1 Fire safety.....	14
4.3.2 Driver's seat configuration.....	14
4.3.3 Minimum passenger capacity	14
4.3.4 Bulkhead	15
4.3.5 Openings (doors, windows, emergency exits)	15
4.3.6 Loading area.....	16
4.4 Patient's compartment.....	18
4.4.1 General.....	18
4.4.2 Safety	18
4.4.3 Hygiene	18
4.4.4 Patient's compartment dimensions	19
4.4.5 Patient and crew seating	24
4.4.6 Ventilation and anaesthetic gas scavenging systems.....	25
4.4.7 Temperature control system	25
4.4.8 Interior lighting	26
4.4.9 Interior noise level	26
4.4.10 Holding system for infusion	26
4.4.11 Retention, fixation and restraint systems.....	27
4.4.12 Mass reserve	27
5 Testing.....	27
5.1 General.....	27
5.2 Testing of the interior noise level.....	27
5.2.1 Specific measurement conditions	27
5.2.2 Measurements.....	28
5.3 Testing of retention systems and fixation of the equipment in the patient's compartment	28
5.3.1 General.....	28
5.3.2 Testing of the stretcher fixation on the vehicle floor.....	30

5.3.3	Testing of the medical devices fixation.....	30
5.3.4	Testing of furniture.....	30
5.3.5	Test procedure	31
5.4	Testing of rounded edges and radius inside the patient's compartment.....	32
5.4.1	Testing of rounded edges.....	32
5.4.2	Testing of radius inside the patient's compartment.....	33
5.5	Procedure to verify the patient's compartment specifications	33
5.6	Procedure to verify the loading area specifications	33
5.6.1	General	33
5.6.2	Procedure to verify the loading angle of 16°	33
5.7	Procedure to verify the dimensions of the patient's compartment	35
5.7.1	Type A and B road ambulances.....	35
5.7.2	Type C road ambulances	35
5.8	Procedure to verify the seats dimensions of the patient's compartment.....	36
5.9	Testing of the ventilation system.....	36
5.10	Testing of the heating system.....	37
5.11	Testing of the cooling system	37
5.11.1	Test procedure	37
5.11.2	Testing of independent air conditioning system	38
5.12	Testing of interior lighting.....	38
5.13	Testing of infusion holding system	38
6	Equipment and medical devices.....	39
6.1	Provision of medical devices	39
6.2	Medical devices storage.....	39
6.3	Requirements for medical devices	39
6.3.1	General	39
6.3.2	Temperature	39
6.3.3	Humidity and ingress of liquids	39
6.3.4	Mechanical strength	39
6.3.5	Fixation of devices.....	40
6.3.6	Electrical safety	40
6.3.7	User interface.....	40
6.3.8	Gas installation	40
6.3.9	Marking and instructions	42
6.3.10	Maintenance	42
6.4	List of equipment.....	42
Annex A (informative)	Test summary.....	51
Annex B (informative)	Recognition.....	52
B.1	Recognition and visibility of ambulances.....	52
B.2	Recognition of crew	52
Annex C (informative)	Hygiene.....	53
Annex D (informative)	A-deviations	54
D.1	Deviation in Spain.....	54
Annex E (normative)	Electric propelled vehicles.....	55
Annex ZA (informative)	Relationship between this European Standard and the essential requirements of Directive 93/42/EEC [O] L 169] aimed to be covered.....	56
Bibliography	57

EN 1789:2020+A1:2023 (E)

European foreword

This document (EN 1789:2020+A1:2023) has been prepared by Technical Committee CEN/TC 239 “Rescue systems”, the secretariat of which is held by DIN.

This European Standard shall be given the status of a national standard, either by publication of an identical text or by endorsement, at the latest by June 2024, and conflicting national standards shall be withdrawn at the latest by June 2024.

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. CEN shall not be held responsible for identifying any or all such patent rights.

A1 This document supersedes EN 1789:2020 **A1**.

This document includes Amendment 1, approved by CEN on 2023-09-21.

The start and finish of text introduced or altered by amendment is indicated in the text by tags **A1** **A1**

This document has been prepared under a standardization request given to CEN by the European Commission and the European Free Trade Association, and supports essential requirements of EU Directive(s).

For relationship with EU Directive(s), see informative Annex ZA which is an integral part of this document.

Any feedback and questions on this document should be directed to the users' national standards body. A complete listing of these bodies can be found on the CEN website.

According to the CEN-CENELEC Internal Regulations, the national standards organisations of the following countries are bound to implement this European Standard: Austria, Belgium, Bulgaria, Croatia, Cyprus, Czech Republic, Denmark, Estonia, Finland, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Malta, Netherlands, Norway, Poland, Portugal, Republic of North Macedonia, Romania, Serbia, Slovakia, Slovenia, Spain, Sweden, Switzerland, Türkiye and the United Kingdom.

Introduction

Road ambulances are subject to a higher risk in use. The exact circumstances of operation cannot always be planned or anticipated in detail.

Vehicles are designed so as to be safe. Design requirements can be derived from European and national occupational safety and health legislation.

Under EU law, employers are responsible for carrying out a risk assessment (89/391/EEC, OSH framework directive) and for provision of safe work equipment (89/655/EEC, use of work equipment directive) that allows employees to work without their health being at risk.

The document was first developed in the late 1990s to define a common approach to requirements to enhance patient and crew safety. The document has evolved and matured through several amendments and revisions.

A1 This revision in 2020 had two key objectives: **A1**

- The first objective was to revise the technical side of the document with more manageable verification in mind, while maintaining the high quality and strict nature of the requirements.
- The second objective was to check all the references and regulations, paying special attention to EU regulations and updated standardization rules.

Testing of special purpose vehicle, such as an ambulance, is complex. Multiple functions (e.g. fixations, maintain systems, noise, illumination, heating, cooling etc.) may require numerous tests, which can be destructive. In this edition, carefully planned tests according to worst-case scenario strategies have reduced the number of destructive tests without sacrificing test qualities.

A1 The previous edition EN 1789:2007+A2:2014 **A1** contained a number of direct references to EU regulations. According to CEN Internal Regulations Part 3:2017 and to avoid duplication as well as outdated references and to enable use of this standard independently of the ECE rules, EU regulations and directives, these references have now been removed from the normative section of the standard.

This document is a reference document which can be used in support of regulations.

For the purpose of verification of an ambulance according to EU vehicle approval process, a section of **A1** EN 1789:2020 **A1** (i.e. patient's compartment) has been referenced directly in Regulation (EU) 2018/858.

A1 The energy sources for motor vehicles are in turmoil due to environmental fight against global climate warming. Alternative energies are becoming more regular in motor vehicles and electric vehicles are already common by most vehicle manufacturers.

In all standardization criteria, combustion engine characteristics have guided the requirements. Therefore, the most obvious ambulance standard requirements (EN 1789) need adjustments introduced by an Amendment to allow verification of electric engine ambulances as compliant to this document. **A1**

EN 1789:2020+A1:2023 (E)

1 Scope

This document specifies requirements for the design, testing, performance and equipping of road ambulances used for the transport, monitoring, treatment and care of patients. It contains requirements for the patient's compartment in terms of the working environment, ergonomic design and the safety of the crew and patients. This document does not cover the training of the crew, which is the responsibility of the authority/authorities in the country where the ambulance is to be registered.

This document is applicable to road ambulances capable of transporting at least one patient on a stretcher and excludes the transportation of hospital beds.

This document also specifies requirements for ambulances intended to carry transport incubator systems.

This document covers the specific requirements of each type of road ambulance, which are designated according to the patient condition.

This document gives general requirements for medical devices carried in road ambulances and used therein and outside hospitals and clinics in situations where the ambient conditions can differ from normal indoor conditions.

2 Normative references

The following documents are referred to in the text in such a way that some or all of their content constitutes requirements of this document. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

Ⓐ₁ EN 16165:2021, *Determination of slip resistance of pedestrian surfaces - Methods of evaluation* Ⓐ₁

EN 3-7:2004+A1:2007, *Portable fire extinguishers - Part 7: Characteristics, performance requirements and test methods*

EN 443:2008, *Helmets for fire fighting in buildings and other structures*

Ⓐ₁ EN 455-1:2020+A1:2022, *Medical gloves for single use - Part 1: Requirements and testing for freedom from holes* Ⓐ₁

EN 455-2:2015, *Medical gloves for single use - Part 2: Requirements and testing for physical properties*

EN 794-3:1998+A2:2009, *Lung ventilators - Part 3: Particular requirements for emergency and transport ventilators*

Ⓐ₁ EN ISO 20417:2021, *Medical devices - Information to be supplied by the manufacturer (ISO 20417:2021, Corrected version 2021-12)* Ⓐ₁

Ⓐ₁ prEN 1865-1:2023,¹ *Patient handling equipment used in ambulances - Part 1: General stretcher systems and patient handling equipment* Ⓐ₁

Ⓐ₁ prEN 1865-2:2022¹, *Patient handling equipment used in ambulances - Part 2: Power assisted stretcher* Ⓐ₁

¹ At draft stage.

prEN 1865-4:2023¹, *Patient handling equipment used in ambulances - Part 4: Foldable patient transfer chair* ^(A1)

prEN 1865-5:2023¹, *Patient handling equipment used in ambulances - Part 5: Stretcher support* ^(A1)

EN 12470-1:2000+A1:2009, *Clinical thermometers - Part 1: Metallic liquid-in-glass thermometers with maximum device*

EN ISO 27427:2019, *Anaesthetic and respiratory equipment - Nebulizing systems and components (ISO 27427:2013)* ^(A1)

EN 13976-1:2018, *Rescue systems - Transportation of incubators - Part 1: Interface requirements*

EN 60601-1:2006+A1:2013, *Medical electrical equipment - Part 1: General requirements for basic safety and essential performance (IEC 60601-1:2005 + Cor.:2006 + Cor.:2007 + A1:2012)*

EN 60601-1-12:2015+A1:2020, *Medical electrical equipment - Part 1-12: General requirements for basic safety and essential performance - Collateral Standard: Requirements for medical electrical equipment and medical electrical systems intended for use in the emergency medical services environment (IEC 60601 1 12:2014 + A1:2020)* ^(A1)

EN 60601-2-4:2011+A1:2019, *Medical electrical equipment - Part 2-4: Particular requirements for the basic safety and essential performance of cardiac defibrillators (IEC 60601 2 4:2010 + A1:2018)* ^(A1)

EN 60601-2-27:2014, *Medical electrical equipment - Part 2-27: Particular requirements for the basic safety and essential performance of electrocardiographic monitoring equipment*

EN ISO 407:2021, *Small medical gas cylinders - Pin-index yoke-type valve connections (ISO 407:2021)* ^(A1)

EN ISO 5359:2014+A1:2017, *Anaesthetic and respiratory equipment - Low-pressure hose assemblies for use with medical gases (ISO 5359:2014 + Amd 1:2017)*

EN ISO 9170-1:2020, *Terminal units for medical gas pipeline systems - Part 1: Terminal units for use with compressed medical gases and vacuum (ISO 9170-1:2017)* ^(A1)

EN ISO 7396-1:2016+A1:2019, *Medical gas pipeline systems - Part 1: Pipeline systems for compressed medical gases and vacuum (ISO 7396-1:2016 + Amd 1:2017)*

EN ISO 10079-1:2022, *Medical suction equipment - Part 1: Electrically powered suction equipment (ISO 10079-1:2022)* ^(A1)

EN ISO 10079-2:2022, *Medical suction equipment - Part 2: Manually powered suction equipment (ISO 10079-2:2022)* ^(A1)

EN ISO 10079-3:2022, *Medical suction equipment - Part 3: Suction equipment powered from a vacuum or positive pressure gas source (ISO 10079-3:2022)* ^(A1)

EN ISO 10524-1:2019, *Pressure regulators for use with medical gases - Part 1: Pressure regulators and pressure regulators with flow-metering devices (ISO 10524-1:2018)*

EN ISO 10524-2:2019, *Pressure regulators for use with medical gases - Part 2: Manifold and line pressure regulators (ISO 10524-2:2018)*

EN 1789:2020+A1:2023 (E)

EN ISO 10524-3:2019, *Pressure regulators for use with medical gases - Part 3: Pressure regulators integrated with cylinder valves (VIPRs) (ISO 10524-3:2019)*

A1 EN ISO 11197:2019, *Medical supply units (ISO 11197:2019)* **A1**

A1 EN ISO 14971:2019+A11:2021, *Medical devices - Application of risk management to medical devices (ISO 14971:2019)* **A1**

A1 prEN ISO 15002:2022¹, *Flow control devices for connection to a medical gas supply system (ISO/DIS 15002:2022)* **A1**

A1 EN ISO 15223-1:2021, *Medical devices - Symbols to be used with information to be supplied by the manufacturer - Part 1: General requirements (ISO 15223-1:2021)* **A1**

EN ISO 19054:2006+A1:2016, *Rail systems for supporting medical equipment (ISO 19054:2005+Amd1:2016)*

EN ISO 20471:2013+A1:2016, *High visibility clothing — Test methods and requirements (ISO 20471:2013, Corrected version 2013-06-01+Amd 1:2016)*

EN ISO 21420:2020-06, *Protective gloves - General requirements and test methods (ISO 21420:2020)*

EN ISO 80601-2-55:2018, *Medical electrical equipment - Part 2-55: Particular requirements for the basic safety and essential performance of respiratory gas monitors (ISO 80601-2-55:2018)*

EN ISO 80601-2-61:2019, *Medical electrical equipment - Part 2-61: Particular requirements for basic safety and essential performance of pulse oximeter equipment (ISO 80601-2-61:2017, Corrected version 2018-02)*

IEC 60364-7-721:2017, *Low-voltage electrical installations — Part 7-721: Requirements for special installations or locations — Electrical installations in caravans and motor caravans*

ISO 3795:1989, *Road vehicles, and tractors and machinery for agriculture and forestry — Determination of burning behaviour of interior materials*

ISO 5128:1980, *Acoustics — Measurement of noise inside motor vehicles*

3 Terms and definitions

For the purposes of this document, the following terms and definitions apply.

ISO and IEC maintain terminological databases for use in standardization at the following addresses:

- IEC Electropedia: available at <http://www.electropedia.org/>
- ISO Online browsing platform: available at <http://www.iso.org/obp>

3.1

patient

person whose condition requires appropriately trained personnel to provide medical care and/or suitable transport

3.2

emergency patient

patient who through sickness, injury or other circumstances is in immediate or imminent danger to life unless emergency treatment and/or monitoring and suitable transport to diagnostic facilities or medical treatment is provided

3.3

road ambulance

vehicle intended to be crewed by a minimum of two appropriately trained crew members for the provision of care and transport of at least one stretchered patient

3.4

type A road ambulance

patient transport ambulance

vehicle designed and equipped for the transport of patients who are not expected to become emergency patients

Note 1 to entry: Two types of patient transport ambulance exist:

- type A₁: suitable for transport of a single patient;
- type A₂: suitable for transport of one or more patient(s) (on stretcher(s) and seat(s)).

3.5

type B road ambulance

emergency ambulance

vehicle designed and equipped for the transport, basic treatment and monitoring of patients

3.6

type C road ambulance

mobile intensive care unit

vehicle designed and equipped for the transport, advanced treatment and monitoring of patients

EN 1789:2020+A1:2023 (E)**3.7****net vehicle mass****<Rescue Service>**

net mass of the road ambulance including the driver taken as 75 kg, 90% fuel tank and all fixed installations

Note 1 to entry: Loose portable patient handling, sanitary, medical and technical equipment are not included in net vehicle mass.

3.8**road ambulance loading capacity**

difference between the permissible gross vehicle mass and the net vehicle mass of the road ambulance

Note 1 to entry: This represents the mass that may be distributed on the road ambulance such that the permissible axle loads are not exceeded.

3.9**fixation system**

system or device to ensure the permanent fixation of medical devices or other equipment into the road ambulance

3.10**retention system**

bracket or other interface device used to secure a mobile or transportable item of equipment or medical device in the road ambulance without the use of tools

3.11**restraint system**

device or combination of devices that minimize movement of the vehicle occupants during crash or major deceleration (e.g. seat belts)

3.12**patient compartment**

interior section of an road ambulance for patient treatment and/or transport

4 Requirements**4.1 General requirements**

Road ambulances and equipment shall, when operated in normal use and maintained according to the instructions of the manufacturer, cause no safety hazard which could reasonably be foreseen using risk management procedures, e.g. in accordance with **EN ISO 14971:2019+A11:2021**, and which is connected with their intended application, in normal condition and in single fault condition.

4.2 Electrical requirements**4.2.1 General**

Electrical installations added to the base vehicle shall comply with those clauses of **IEC 60364-7-717:2009** which are applicable to road ambulances. For the supply system of the medical equipment **EN 60601-1:2006+A1:2013** and **EN 60601-1-12:2015+A1:2020**, Clause 11 shall apply.

4.2.2 Electromagnetic compatibility (EMC)

In order to minimize the risk to the safe operation of the complete road ambulance and any of the equipment operated on or in the vehicle from the effects of electromagnetic influences:

- Communication equipment (e.g. radio installation) shall comply with national and/or European regulations.
- The complete operational vehicle shall consist of components, equipment or sub systems that comply or are certified as conforming to the respective industry EMC regulations.

A road ambulance as supplied and certified may not be fully equipped and therefore some responsibility for added equipment after conversion rests with the customer/user.

4.2.3 Battery and alternator

Batteries shall be positioned to allow maintenance without removing the battery from its securing device. The construction of the battery and all connections to it shall be such as to prevent any possibility of an inadvertent short circuit.

For types A2, B and C road ambulances the electrical system shall be capable of holding a reserve of electrical power for restarting the engine. The characteristics of the alternator, the starter batteries as well as additional batteries, if fitted, shall comply with Table 1.

Additional batteries may be required to power the medical devices carried on board and the intended use of the road ambulance.

Table 1 — Minimum capacity/power

		Type of road ambulance			
		A ₁	A ₂	B	C
Starter battery(ies)	Nominal voltage 12 V	54 Ah	54 Ah up to 4 seats and 80 Ah for more than 4 seats in the patient compartment	80 Ah	80 Ah
	Nominal voltage 24 V	–	–	63 Ah (2 × 12 V)	63 Ah (2 × 12 V)
Additional battery(ies) ^b	Nominal voltage 12 V	–	–	80 Ah ^a	80 Ah
	Nominal voltage 24 V	–	–	63 Ah ^a (2 × 12 V)	63 Ah (2 × 12 V)
Alternator power		700 W	700 W	1 200 W	1 200 W
<p>^a Recommended for special operational conditions.</p> <p>^b Additional batteries shall have high cyclic stability (e.g. gel batteries) and shall be of a sealed type.</p>					

When the engine is idling electrical stability should be maintained between electrical load and alternator output. In order to achieve this it may be necessary to fit an electrical load prioritization device to the vehicle.

EN 1789:2020+A1:2023 (E)

4.2.4 Electrical installation

4.2.4.1 In type B and C road ambulances there shall be a recessed externally mounted power connector to enable external power to be provided for operations such as the following:

- charging battery(ies);
- operating medical devices, when installed;
- operating a patient compartment heater, when installed;
- operating an engine preheater, when installed.

The connector for 110 V or 220/240 V shall be a male connector and not interfere with the electrical and mechanical safety.

It shall not be possible to start the engine whilst it is connected to an external power supply unless an automatic mechanical disconnection is fitted.

If no automatic mechanical disconnection is fitted, the connector shall be on the driver's side.

The 110 V or 220/240 V circuit shall be protected either by an "earth leakage device" with a maximum setting of 30 mA or by a separate transformer. If the protection is given only by an "earth leakage device" there shall be a label near the plug that reads as follows: "CAUTION! CONNECT ONLY TO AN AUTHORISED SOCKET."

4.2.4.2 A minimum number of separately protected 12 V DC outlets shall be available according to Table 2. The outlets shall be available for medical devices, located in the area of use and in the storage area. The nominal voltage shall be 13,8 V. Voltage fluctuations shall not exceed the range of 12,4 V and 15,1 V.

The power supply shall continuously supply the medical devices with electrical power with the engine running. The outlets for the medical devices shall be labelled with the nominal voltage and current rating. The outlets shall have a visible indication under intended operational conditions in order to show if there is voltage on the outlet.

If the road ambulance is intended to carry a transport incubator system it shall have a four-pole connector as specified in EN 13976-1:2018, subclause 4.2.3, Figure 2. In that case the nominal current needs to be assessed (minimum 23 Amp according to EN 13976-1:2018, subclause 4.1.3).

Table 2 — 12 V connections for medical devices in patient's compartment

	Type of road ambulance							
	A ₁		A ₂		B		C	
Minimum number of connections	1	1	1	1	3	1	3	1
Minimum capacity in Ampere	10	15	10	15	10	15	10	15

4.2.4.3 Any additional electrical systems fitted to the base vehicle shall be separate from the base vehicle electrical system and the body or chassis shall not be used as an earth return for additional circuits.