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Space systems — Fluid characteristics, sampling and test methods —

Part 4: **Helium**

Systèmes spatiaux — Caractéristiques, échantillonnage et méthodes d'essai des fluides —

Partie 4: Hélium



Reference number ISO 15859-4:2004(E)

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Foreword

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International Standards are drafted in accordance with the rules given in the ISO/IEC Directives, Part 2.

The main task of technical committees is to prepare International Standards. Draft International Standards adopted by the technical committees are circulated to the member bodies for voting. Publication as an International Standard requires approval by at least 75 % of the member bodies casting a vote.

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

ISO 15859-4 was prepared by Technical Committee ISO/TC 20, Aircraft and space vehicles, Subcommittee SC 14, Space systems and operations.

ISO 15859 consists of the following parts, under the general title *Space systems* — *Fluid characteristics*, *sampling and test methods*:

— Part 1: Oxygen— Part 2: Hydrogen— Part 3: Nitrogen

– ∃Part 4: Helium

- Part 5: Nitrogen tetroxide propellants
- Part 6: Monomethylhydrazine propellant
- Part 7: Hydrazine propellant
- Part 8: Kerosine propellant
- Part 9: Argon
- Part 10: Water
- Part 11: Ammonia
- Part 12: Carbon dioxide
- Part 13: Breathing air

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Introduction

Fluid operations at a spaceport or launch site may involve a number of operators and supplier/customer interfaces, from the fluid production plant to the delivery to the launch vehicle or spacecraft. The purpose of ISO 15859 is to establish uniform requirements for the components, sampling and test methods of fluids used in the servicing of launch vehicles, spacecraft and ground support equipment. The fluid composition limits specified are intended to define the purity and impurity limits of the fluid for loading into the launch vehicle or spacecraft. The fluid sampling and test methods are intended to be applied by any operator. The fluid sampling and test methods are acceptable methods for verification of the fluid composition limits.

Space systems — Fluid characteristics, sampling and test methods —

Part 4: **Helium**

1 Scope

This part of ISO 15859 specifies limits for the composition of helium and establishes the sampling and test requirements applicable for the verification of the helium composition.

This part of ISO 15859 is applicable to helium, used in both flight hardware and ground facilities, systems and equipment, of the following types and grades.

- Type I: gaseous
 - Grade A: purging and pressurizing helium;
 - Grade F: purging and pressurizing helium;
 - Grade J: purging and pressurizing helium;
- Type II: liquid
 - Grade A: purging and pressurizing helium;
 - Grade F: purging and pressurizing helium.

This part of ISO 15859 is applicable to influents only within the specified limits herein.

This part of ISO 15859 is applicable to any sampling operation required to ensure that, when the fluid enters the launch vehicle or spacecraft, the fluid composition complies with the limits provided hereafter or with any technical specification agreed to for a particular use.

2 Normative references

The following referenced documents are indispensable for the application 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.

ISO 9000, Quality management systems — Fundamentals and vocabulary

Terms and definitions

For the purposes of this document, the terms and definitions given in ISO 9000 and the following apply.

3.1

total hydrocarbon (as methane)

the single carbon atom equivalent

3.2

verification test

analysis performed on the fluid in the container, or a sample thereof, which is representative of the supply, permitting the verification of fluid composition limits

Chemical composition 4

Unless otherwise provided in an applicable technical specification, the composition of helium delivered to the flight vehicle interface shall be in accordance with the limits given in Table 1 when tested in accordance with the applicable test methods.

Table 1 — Composition limits

Component			Limits					
			Type I (gaseous)			Type II (liquid)		
			Grade A	Grade F	Grade J	Grade A	Grade F	
Purity	Helium	olume fraction, %, min.	99,99	99,995	99,999	99,99	99,995	
Impurities	Water	μl/l, max.	9	5	1,9	9	3	
	Hydrocarbons (as methane)	μl/l, max.	5	10	0,1	5	1	
	Oxygen	μl/l, max.	10	5	1	10	3	
	Nitrogen	μl/l, max.	50	20	3	50	5	
	Neon	μl/l, max.	_	23	_	_	23	
	Argon	μl/l, max.	_	_	_	_	1	
	Hydrogen	μl/l, max.	_	_	_	_	5	
	Carbon monoxide plus carbon dioxide	μl/l, max.		1	0,1	_	1	
	Total allowable impurit	es μl/l, max.	100	50	10	100	50	

Procurement

Helium types and grades specified in Clause 1 should be procured in accordance with an applicable national standard.

6 Fluid sampling

CAUTION — Gaseous helium is an asphyxiant. Human contact with liquid helium will result in severe injury. Care should be taken in the handling and storage of liquid helium to prevent contact with the human body. Care should also be taken to prevent high concentrations of gaseous helium in confined spaces.

6.1 Plan

In order to ensure that the fluid composition complies with the limits specified in this part of ISO 15859, a fluid sampling plan should be established by all the involved operators, from the production to the space vehicle interface, and approved by the final user. Sampling activities and test methods shall comply with all safety regulations and rules applicable to that task. This plan shall specify

—	the sampling points,
_	the sampling procedures,
_	the sampling frequency,
	the sample size,
_	the number of samples,

the responsibilities of any involved operator.

6.2 Responsibility for sampling

Unless otherwise provided in an applicable technical specification, the helium delivered to the flight vehicle interface shall be sampled and verified by the supplier responsible for providing the helium to the flight vehicle. The supplier may use his/her or any other resources suitable for the performance of the verification tests specified herein unless otherwise directed by the customer.

6.3 Sampling points

— the test methods, and

Unless otherwise specified, sampling shall be conducted at the fluid storage site or the flight vehicle interface.

6.4 Sampling frequency

Sampling shall be performed annually or in accordance with a time agreed upon by the supplier and the customer.

6.5 Sample size

The quantity in a single sample container shall be sufficient to perform the analysis for the limiting characteristics. If a single sample does not contain a sufficient quantity to perform all of the analyses for the required quality verification test, additional samples shall be taken under similar conditions.

6.6 Number of samples

The number of samples shall be in accordance with one of the following:

- a) one sample per storage container;
- b) any number of samples agreed upon by the supplier and the customer.

6.7 Storage container

Unless otherwise provided by the applicable sampling plan, the fluid storage container shall not be refilled after the sample is taken.

6.8 Gaseous samples

Gaseous samples shall be a typical specimen from the gaseous supply. Samples shall be obtained in accordance with one of the following.

- a) By filling the sample container and storage containers at the same time, on the same manifold, under the same conditions and with the same procedure.
- b) By withdrawing a sample from the supply container through a suitable connection into the sample container. No pressure regulator shall be used between the supply and the sample containers. (Suitable valves are permissible.) For safety reasons, the sample container and sampling system shall have a rated service pressure at least equal to the pressure in the supply container.
- c) By connecting the container being sampled directly to the analytical equipment using suitable pressure regulation to prevent over-pressurizing this equipment.

6.9 Liquid samples (vaporized)

Vaporized liquid samples shall be a typical specimen from the liquid supply. Samples shall be obtained by flowing liquid from the supply container into or through a suitable container in which a representative liquid sample is collected and then completely vaporized.

6.10 Rejection

When any sample of the fluid tested in accordance with Clause 7 of this part of ISO 15859 fails to conform to the requirements specified herein, the fluid represented by the sample shall be rejected. Disposal of the rejected fluid shall be specified by the customer.

7 Test methods

7.1 General

The supplier will ensure, by standard practice, the quality level of helium. If required, alternate test methods are described in 7.3 to 7.10. Other test methods not listed in this part of ISO 15859 are acceptable if agreed upon between the supplier and the customer.

These tests are a single analysis or a series of analyses performed on the fluid to ensure the reliability of the storage facility to supply the required quality level. This can be verified by analysis of representative samples of the fluid from the facility at appropriate intervals as agreed upon between the supplier and the customer. Tests may be performed by the supplier or by a laboratory agreed upon between the supplier and the customer.

The analytical requirements for the tests shall include the determination of all limiting characteristics of helium.

7.2 Parameters of analysis

The parameters for analytical techniques contained in 7.3 to 7.10 are the following:

a) purity and impurity contents shall be expressed as a percentage by volume (volume fraction, %) unless otherwise noted:

- b) calibration gas standards containing the applicable gaseous components may be required to calibrate the analytical instruments used to determine the limiting characteristic levels of fluid;
- c) if required by the customer, the accuracy of the measuring equipment used in preparing these standards shall be traceable to an established institute for standards;
- d) analytical equipment shall be operated in accordance with the manufacturer's instructions.

7.3 Helium purity

The purity of helium shall be determined by one of the following procedures.

- a) By a thermal conductivity analyser measuring the aggregate impurities which have different thermal conductivities than helium. The analyser is to be calibrated at appropriate intervals using calibration gas standards. The range of the analyser shall be no greater than 10 times the difference between the specified minimum value of helium purity, expressed as a volume fraction (%), and 100 %.
- b) By determining the quantity of the aggregate impurities by a mass spectrometer. The purity of helium is the value obtained when the quantity of aggregate impurities, expressed as a volume fraction (%), is subtracted from 100.
- c) By determining the quantity of aggregate impurities using the methods in 7.4 to 7.10. The purity of helium is the value obtained when the quantity of aggregate impurities, expressed as a volume fraction (%), is subtracted from 100.
- d) By gas chromatography system in accordance with 7.7 a) using a carrier gas other than helium.

7.4 Water content

For liquid helium, the water content cannot be determined by sampling. For gaseous helium, the water content shall be determined by one of the following procedures.

- a) By a dew-point analyser in which the temperature of a viewed surface is measured at the time water first begins to form.
- b) By a piezoelectric sorption hygrometer, of which the accuracy of analysis shall be \pm 0,1 cm³/m³ or 5 % of the reading, whichever is greater.
- c) By a metal-oxide-capacitor-equipped analyser within a range which is no greater than 10 times the specific maximum water content.
- d) By an electrolytic hygrometer having an indicator graduated in cubic centimetres per cubic metre within a range which is not greater than 10 times the specified maximum water content.

7.5 Total hydrocarbon content (THC)

The total (volatile) hydrocarbon content (as methane) shall be determined by one of the following procedures.

- a) By a gas chromatograph method such as described under 7.7 a).
- b) By a gas-cell-equipped infrared analyser. The analyser shall be calibrated at appropriate intervals by use of calibration gas standards at a wavelength of approximately 3,5 µm (the characteristic absorption wavelength for C-H stretching). The analyser shall be operated so that its sensitivity for methane is at least 10 % of the specified maximum total hydrocarbon contents.

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Oxygen content 7.6

The oxygen content shall be determined by one of the following procedures.

- a) By a gas chromatography method such as that described under 7.7 a).
- b) By a mass spectrometer. The mass spectrometer shall be operated so that its sensitivity is at least 10 % of the specified oxygen content.
- By an electrochemical-type oxygen analyser containing a solid or an aqueous electrolyte. The analyser shall be calibrated at appropriate intervals by use of calibration gas standards or integrally in accordance with Faraday's Law. The range used should be no greater than 10 times the specified maximum oxygen content.
- d) By a heat-of-reaction-type analyser. The analyser shall be calibrated at appropriate intervals by the use of calibration gas standards or integrally in accordance with Faraday's Law. The range used should be no greater than 10 times the specified maximum oxygen content.
- By an analyser in which oxygen reacts to form a compound which is subsequently measured. The analyser shall be calibrated at appropriate intervals by the use of calibration standards. The range used shall be no greater than 10 times the specified maximum oxygen content.

Argon, neon, and nitrogen content

The argon, neon, and nitrogen contents shall be determined by one of the following procedures.

- By a gas chromatograph. This method may be used not only for argon, neon and nitrogen determination but also for the determination of any other limiting characteristic gaseous components (see Annex A). The analyser shall be capable of separating and detecting the component with a sensitivity of 10 % of the specified maximum amount of the component. Appropriate impurity concentrating techniques may be used to attain the sensitivity. The analyser shall be calibrated at appropriate intervals by the use of calibration gas standards.
- b) By a mass spectrometer. The mass spectrometer shall be operated so that its sensitivity is at least 10 % of the specified maximum amount of the component.

Hydrogen content 7.8

The hydrogen content shall be determined by one of the following procedures.

- a) By a gas chromatograph in accordance with 7.7 a).
- By a mass spectrometer. The mass spectrometer shall be operated so that its sensitivity is at least 10 % of the specified hydrogen content.
- By an analyser in which hydrogen reacts to form a compound which is subsequently measured. The analyser shall be calibrated at appropriate intervals by the use of calibration gas standards. The range used shall be no greater than 10 times the specified maximum hydrogen content.

7.9 Carbon dioxide content

The carbon dioxide content shall be determined by one of the following procedures.

- a) By a gas chromatography method such as that described in 7.7 a). The technique utilized shall be specific for the separation and analysis of carbon dioxide.
- b) By a mass spectrometer. The mass spectrometer shall be operated so that its sensitivity is at least 10 % of the specified maximum amount of the component.

- c) By a catalytic methanizer gas chromatography method such as that described in 7.7 a).
- d) By an analyser in which carbon dioxide reacts to form a compound which is subsequently measured. The analyser shall be calibrated at appropriate intervals by the use of calibration standards. The range used shall be no greater than 10 times the specified maximum carbon dioxide content.

7.10 Carbon monoxide content

The carbon monoxide content shall be determined by one of the following procedures.

- a) By a gas chromatography method such as that described under 7.7 a). The technique utilized shall be specific for separation and analysis of carbon monoxide.
- b) By an analyser in which carbon monoxide reacts to form a compound which is subsequently measured. The analyser shall be calibrated at appropriate intervals by the use of calibration standards. The range used shall be no greater than 10 times the specified maximum carbon monoxide content.
- c) By a catalytic methanizer gas chromatography method such as that described under 7.7 a).

Annex A

(informative)

Gas chromatography (GC) and mass spectrometer (MS) applications

Gas chromatography (GC) should be used as the reference or preferred method to analyse helium impurities, except for water content.

A gas chromatograph coupled with a mass spectrometer (GC-MS) may be used as an alternative to simple gas chromatography to avoid possible interference (especially for the hydrocarbons).

Table A.1 summarizes the applications of these methods for helium.

Table A.1 — Application of GC and MS

Component	GC with DID detector on molecular sieve column	GC with FID detector on Porapak ^a column (or equivalent)	GC with methanizer and FID detector on Porapak ^a column (or equivalent)	GC-MS	MS
Water	_	_	_	_	_
Hydrocarbon (as methane)	Х	_	_	Х	_
Oxygen	_	X	X	_	_
Argon, neon, nitrogen	X	_	_	Х	Х
Hydrogen	_	X	Х	_	_
Carbon dioxide			X	Х	_
Carbon monoxide	Х	_	Х	Х	_

DID = Discharge ionization detector

FID = Flame ionization detector

[&]quot;X" indicates that the method can be used.

^{-&}quot; indicates that the method is not used.

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