

**ANSI/CGA M-1—2018
STANDARD FOR MEDICAL
GAS SUPPLY SYSTEMS AT
HEALTH CARE FACILITIES**

FOURTH EDITION



Currently in preview, click buy full version

PLEASE NOTE:

The information contained in this document was obtained from sources believed to be reliable and is based on technical information and experience currently available from members of the Compressed Gas Association, Inc. and others. However, the Association or its members, jointly or severally, make no guarantee of the results and assume no liability or responsibility in connection with the information or suggestions herein contained. Moreover, it should not be assumed that every acceptable commodity grade, test or safety procedure or method, precaution, equipment or device is contained within, or that abnormal or unusual circumstances may not warrant or suggest further requirements or additional procedure.

This document is subject to periodic review, and users are cautioned to obtain the latest edition. The Association invites comments and suggestions for consideration. In connection with such review, any such comments or suggestions will be fully reviewed by the Association after giving the party, upon request, a reasonable opportunity to be heard. Proposed changes may be submitted via the Internet at our website, www.cganet.com.

This document should not be confused with federal, state, provincial, or municipal specifications or regulations; insurance requirements; or national safety codes. While the Association recommends reference to or use of this document by government agencies and others, this document is purely voluntary and not binding unless adopted by reference in regulations.

A listing of all publications, audiovisual programs, safety and technical bulletins, and safety posters is available via the Internet at our website at www.cganet.com. For more information contact CGA. Phone: 703-788-2700, ext. 799. E-mail: customerservice@cganet.com.

Work Item 18-027
Medical Gases Committee

NOTE—Technical changes from the previous edition are underlined.

NOTE—Appendices A, B, and C (Informative) are for informational only.

FOURTH EDITION: 2018
THIRD EDITION: 2013
SECOND EDITION: 2007
FIRST EDITION: 2003

© 2018 The Compressed Gas Association, Inc. All rights reserved.

All materials contained in this work are protected by United States and international copyright laws. No part of this work may be reproduced or transmitted in any form or by any means, electronic or mechanical including photocopying, recording, or any information storage and retrieval system without permission in writing from The Compressed Gas Association, Inc. All requests for permission to reproduce material from this work should be directed to The Compressed Gas Association, Inc., 8484 Westpark Drive, Suite 220, McLean, VA 22102. You may not alter or remove any trademark, copyright or other notice from this work.

AMERICAN NATIONAL STANDARD

Approval of an American National Standard requires verification by ANSI that the requirements for due process, consensus, and other criteria for approval have been met by the standards developer.

Consensus is established when, in the judgment of the ANSI Board of Standards Review, substantial agreement has been reached by directly and materially affected interests. Substantial agreement means much more than a simple majority, but not necessarily unanimity. Consensus requires that all views and objections be considered, and that a concerted effort be made toward their resolution.

The use of American National Standards is completely voluntary; their existence does not in any respect preclude anyone, whether he has approved the standards or not, from manufacturing, marketing, purchasing, or using products, processes, or procedures not conforming to the standards.

The American National Standards Institute does not develop standards and will in no circumstances give an interpretation of any American National Standard. Moreover, no person shall have the right or authority to issue an interpretation of an American National Standard in the name of the American National Standards Institute. Requests for interpretations should be addressed to the secretariat or sponsor whose name appears on the title page of this standard.

CAUTION NOTICE: The American National Standard may be revised or withdrawn at any time. The procedures of the American National Standards Institute require that action be taken to periodically reaffirm, revise, or withdraw this standard. Purchasers of American National Standards may receive current information on all standards by calling or writing the American National Standards Institute, 11 West 42nd Street, New York, NY 10036.

Approved as an American National Standard on April 6, 2018.

Contents	Page
1 Introduction.....	1
2 Scope	1
3 Definitions.....	2
4 Health hazards and safety considerations	5
5 Applicable codes and standards	6
6 Personnel	6
6.1 Quality control unit.....	6
6.2 Personnel qualifications.....	7
7 Site selection	9
7.1 Meeting code setback distances	9
7.2 Health care facility, system owner, and supplier responsibilities.....	10
8 Developing a suitable site	11
8.1 Equipment layout.....	11
8.2 Foundations.....	11
8.3 Security.....	11
9 Equipment selection	12
9.1 System specifications	12
9.2 Equipment specifications.....	12
9.3 System design	15
9.4 Process flow diagrams	17
9.5 Regulator set points.....	17
9.6 Line pressure relief valves (thermal reliefs).....	18
9.7 Vendor selection.....	18
9.8 Equipment reuse	18
10 Equipment transportation	18
11 Equipment installation	18
11.1 Temporary system.....	18
11.2 Cleaning	19
11.3 Odor test.....	20
11.4 Installation	20
11.5 Pipe joints.....	20
11.6 System identification, marking, and tagging.....	22
11.7 System pressure test.....	22
12 Startup.....	22
12.1 First fill.....	22
12.2 Purge	23
12.3 Product testing.....	23
12.4 Startup testing	23
12.5 System verification	23
12.6 Commissioning.....	24
13 Operation.....	25
13.1 Maintenance.....	25
13.2 Reporting complaints.....	26
13.3 Product recall.....	26
14 System removal.....	26

15 Documentation 26

 15.1 Installation/modification 26

 15.2 Equipment inspection 27

16 References 28

17 Additional information 29

Appendices

Appendix A—Ventilation [3, 4] (Informative) 30

Appendix B—Switching vaporizers for medical gas supply systems [3, 4] (Informative) 31

Appendix C—NPFA 55 and NPFA 99 requirements for bulk medical gas supply systems [3, 4] (Informative) 32

Currently in preview, click buy full version

1 Introduction

Compressed medical gases (CMG) are intended for use in the diagnosis, cure, mitigation, treatment, or prevention of disease in human beings or animals.

Medical gas supply systems deliver CMG to piped distribution systems at health care facilities. Oxygen USP and medical air USP provide direct assistance for breathing and also assist in the delivery of medical treatment(s). Oxygen USP, medical air USP, and CMG mixtures are used in hyperbaric chambers. Nitrogen NF is used to operate power tools during patient care procedures. CMG shall be permitted to be supplied to the medical gas piped distribution system by cylinders or containers connected to a manifold system, a cryogenic fluid central supply (either a permanently installed bulk supply system or permanently installed microbulk supply system), or a temporary supply system.

Supplier-owned or health care facility-owned storage and control systems that supply CMG shall be installed and maintained in compliance with this standard, applicable regulations, and the U.S. Food and Drug Administration (FDA) current good manufacturing practices (CGMP).

The system owner is responsible for compliance with CGMP. Title 21 of the U.S. Code of Federal Regulations (21 CFR) Parts 210-211 prohibits the adulteration of drugs [1].¹ Introducing a drug product into a system that has not been installed in accordance with CGMP will adulterate the product. The supplier of the CMG is responsible for ensuring that the system has been installed in compliance with CGMP prior to first filling of the system. Supplier and contractor personnel that work on supplier-owned equipment shall be able to provide documentation that demonstrates compliance with this standard.

2 Scope

This standard provides the minimum requirements for the design, installation, maintenance, testing, and removal of CMG supply systems at health care facilities. For facilities that are solely intended for use in non-human applications (i.e., veterinary or pharmaceutical), the applicability of this standard is to be determined by the CMG system designer, authority having jurisdiction (AHJ), or other related parties based on facility requirements. Strict adherence to CGMP shall be taken into account to prevent adulteration of the CMG.

This standard applies to all new or upgraded CMG supply systems at health care facilities. It provides direction for compliance with the following national regulations and model codes:

- Federal *Food, Drug, and Cosmetic Act* [2];
- Title 21 of the U.S. *Code of Federal Regulations (21 CFR) Parts 210 to 211* [1];
- NFPA 55, *Compressed Gases and Cryogenic Fluids Code* [3]; and
- NFPA 99, *Health Care Facilities Code* [4].

Section 5 covers the scope of these regulations and their applicability to CMG supply systems. This standard captures the requirements from these codes along with best practices to provide a comprehensive publication for the process of designing, locating, installing, commissioning, maintaining, testing, removing, and documenting work on a medical gas supply system.

In an upgrade, the complete system shall be modified to comply with current standards. Failure of individual components may not invoke a full system upgrade. An exception is replacing piping or control assemblies that do not include the replacement or relocation of the bulk liquid storage vessel(s) may not invoke the foundation requirements in 8.2.

This standard does not apply to:

- piped distribution systems including source valve requirements;

¹ References are shown by bracketed numbers and are listed in the order of appearance in the reference section.

- manufacture of assemblies such as pressure control manifolds that are manufactured in a supplier's shop and qualified for medical gas service in accordance with the CMG supply system installer's policies;
- manufacturing plants or other establishments operated by the supplier or the supplier's agent for the purpose of storing and refilling portable containers, trailers, mobile supply trucks, or tank cars with medical gases;
- nonhealth care facilities such as laboratories, pharmaceutical, or biotechnology facilities;
- medical vacuum systems; or
- supply systems that generate CMG on-site.

3 Definitions

For the purpose of this standard, the following definitions apply.

3.1 Publication terminology

3.1.1 Shall

Indicates that the procedure is mandatory. It is used wherever the criterion for conformance to specific recommendations allows no deviation.

3.1.2 Should

Indicates that a procedure is recommended.

3.1.3 May

Indicates that the procedure is optional.

3.1.4 Will

Is used only to indicate the future, not a degree of requirement.

3.1.5 Can

Indicates a possibility or ability.

3.2 Technical definitions

3.2.1 Argon

Inert gas used as a purge medium during the fabrication of welded stainless steel piping assemblies.

3.2.2 Authority having jurisdiction (AHJ)

Organization, office, or individual responsible for enforcing the requirements of a code or standard or responsible for approving equipment, materials, installations, or procedures.

NOTE—There may be multiple AHJs with various levels of responsibilities and authority.

3.2.3 Brazer

Person with documented proficiency in the silver brazing procedures used to join pipe and tubing during the installation of a CMG supply system.

3.2.4 Bulk cryogenic fluid central supply system

Cryogenic fluid central supply system with a storage capacity greater than 566 m³ (20 000 scf).

3.2.5 Certified

Container or system whose content is manufactured, tested, and in compliance with applicable standards, for example, *United States Pharmacopoeia–National Formulary (USP–NF)* or supplier standards [5].

3.2.6 Clean for oxygen service

System or component of a system that is specifically cleaned, inspected, labeled, and packaged for use with oxygen.