LEARNING OBJECTIVES

2.07.01 Explain the purpose of respiratory protection standards and regulations.

2.07.02 Identify the OSHA, ANSI, and DOE respiratory protection program requirements.

2.07.03 Identify the standards which regulate respiratory protection.

2.07.04 Describe the advantages and disadvantages (limitations) of each of the following respirators:
   a. Air purifying, particulate removing filter respirators
   b. Air purifying, Chemical Cartridge and Canister respirators for Gases and Vapors
   c. Full-face, supplied-air respirator
   d. Self-contained breathing apparatus (SCBA)
   e. Combination atmosphere supplying respirators.

2.07.05 Define the term protection factor (PF).

2.07.06 State the difference between a qualitative and quantitative fit test.

2.07.07 State the recommended physical functions the subject must perform during a respirator fit test.

2.07.08 State how the term protection factor (PF) is applied to selection of respiratory protection equipment.

2.07.09 State the general considerations and considerations for the nature of the hazard when selecting the proper respiratory protection equipment.

2.07.10 Identify the types of respiratory equipment available for use at your site.

2.07.11 Identify the quality specifications breathing air must meet.
2.07 - RESPIRATORY PROTECTION

2.07.01 Explain the purpose of respiratory protection standards and regulations.

2.07.02 Identify the OSHA, ANSI, and DOE respiratory protection program requirements.

2.07.03 Identify the standards which regulate respiratory protection.

OSHA AND DOE REQUIREMENTS

The Occupational Safety and Health Standard, 29 CFR, Part 1910.134, specifies the minimal acceptable respiratory protection program must contain or address the following:

- Written standard operating procedures governing the selection and use of respirators shall be established.
- Respirators shall be selected on the basis of hazards to which the worker is exposed.
- The user shall be instructed and trained in the proper use of respirators and their limitations.
- Where practicable, the respirators should be assigned to individual workers for their exclusive use.
- Respirators shall be regularly cleaned and disinfected. Those issued for the exclusive use of one worker should be cleaned after each day's use, or more often if necessary. Those used by more than one worker shall be thoroughly cleaned and disinfected after each use.
- Respirators shall be stored in a convenient, clean, and sanitary location.
- Respirators used routinely shall be inspected during cleaning. Worn or deteriorated parts shall be replaced. Respirators for emergency use such as self-contained devices shall be thoroughly inspected at least once a month and after each use.
- Appropriate surveillance of worker area conditions and degree of employee exposure or stress shall be maintained.
There shall be regular inspection and evaluation to determine the continued effectiveness of the program.

Persons should not be assigned to tasks requiring use of respirators unless it has been determined that they are physically able to perform the work and use the equipment. The local physician shall determine what health and physical conditions are pertinent. The respirator user's medical status should be reviewed periodically (for instance, annually).

Approved or accepted respirators shall be used when they are available. The respirator furnished shall provide adequate respiratory protection against the particular hazard for which it is designed in accordance with standards established by competent authorities.

These "Eleven Commandments" form the basis for any occupational safety respiratory protection program. ANSI Z88.2-1980 further specifies the minimal acceptable program for industries involved in the use of radioactive material.

If allowance for the use of respiratory protection equipment in estimating exposures is made, then the following must be observed:

- The protection factor for the device selected must be greater than the ratio of the peak exposure concentration and the associated DAC.

- The average concentration inhaled on any one day must be less than the associated DAC.

- If the exposure is later found to be greater than estimated, the corrected value shall be used, if less than estimated, the corrected value may be used.

- Surveys and bioassays conducted as appropriate to evaluate actual exposures.

- Written procedures for selection, fitting, maintenance, records, issuance and pre-use operability checks of respirators, and supervision and training of personnel using respirators must be established.

- Prior to initial use and annually, determination by a physician of a user's physical capability to wear a respirator must be performed.

- A written policy statement on use of engineering controls instead of respirators; routine, non-routine, and emergency use of respirators; and periods of respirator
use and relief from respirator use must be issued.

- Each user must be advised that they can leave the work area upon failure of equipment, physical distress or deterioration of operating conditions.

- Equipment is to be used for appropriate environment and special equipment such as visual or communication devices are to be issued when needed.

- Emergency use equipment must be specifically certified as such by NIOSH/MSHA.

DOE RCM Respiratory Protection Program Requirements

1. Use of respiratory protection shall be reduced to the minimum practicable by implementing engineering controls and work practices to contain radioactivity at the source.

2. DOE 5480.4 mandates the requirements contained in ANSI Z88.2 and 29 CFR 1910.134 for implementation of the Respiratory Protection Program and associated training of personnel.

3. Respirators shall be issued only to personnel who are trained, fitted and medically qualified to wear the specific type of respirator. Training and qualification testing shall be performed annually.

4. Positive controls shall be maintained for the issue, use and return of respirators to ensure that only qualified personnel wear respirators.

5. DOE 5480.4 mandates that breathing air meet the specifications of ANSI/CGA G-7.1 Grade D breathing air as specified in 29 CFR 1910.134. Compressed air supplied to respirators shall be tested quarterly. Compressors shall be of the breathing air type and shall not allow oil or other chemicals and fumes to enter the breathing air supply. Special attention shall be focused on the location of compressor air supply intakes and on cross-connections to other compressed gas systems to prevent contamination.

6. Facility safety analyses should not take credit for the use of respiratory protection for routine work involving potential exposure to airborne radioactive materials. Engineering controls should be designed to control radioactive materials at the source, so that respiratory protection can be reduced.
RESPIRATORY PROTECTION EQUIPMENT

2.07.04 Describe the advantages and disadvantages (limitations) of each of the following respirators:

a. Air purifying, particulate removing filter respirators

Air Purifying, Particulate-Removing Filter Respirators

Description:

These are generally called "dust," "mist," or "fume" respirators and by a filtering action remove particulates before they can be inhaled. Single use, quarter mask, half mask, full facepiece, and air powered hood/mask are the five types of respirators that work by the particulate removal method. Air purifying respirators generally operate in the negative pressure (NP) mode; that is, a negative pressure is created in the facepiece during inhalation. An exception is a special type of powered air purifying respirator that operates continuously in the positive pressure (PP) mode by using a motor-driven blower to drive the contaminated air through an air purifying filter or sorbent canister. In power reactors only the half face, full-piece, and air powered hood are respirators approved for use in protection against airborne radionuclides.

Limitations:

Air purifying respirators do not provide oxygen, so they must NEVER be worn in oxygen-deficient atmospheres.

Particulate-removing air-purifying respirators offer no protection against atmospheres containing contaminant gases or vapors.

Except for pressurized air purifier respirators, these respirator types should not be used for abrasive blasting operations.

Battery operated air powered respirators are limited by battery life which may be unknowingly shortened due to a memory build-up on the rechargeable NiCd batteries.

High humidity may increase breathing resistance as paper elements become water saturated.
2.07.04 Describe the advantages and disadvantages (limitations) of each of the following respirators:

b. Air purifying, Chemical Cartridge and Canister respirators for Gases and Vapors

Air Purifying, Chemical Cartridge and Canister Respirators for Gases and Vapors

Description:

Vapor and gas-removing respirators use cartridges or canisters containing chemicals (i.e., sorbents) to trap or react with specific vapors and gases and remove them from the air breathed. The basic difference between a cartridge and a canister is the volume of the sorbent. In power reactors, a combination filter-canister type respirators are used with tight-fitting full facepiece respirators and are used for protection against radioiodines.

Limitations:

These respirators do not provide oxygen, so they must NEVER be worn in oxygen deficient atmospheres.

Unless specifically approved by DOE, no credit may be taken for the use of sorbent cartridges or canisters for protection against radioactive gases and vapors.

High humidity environments will shorten the life of the sorbent material.
2.07.04 Describe the advantages and disadvantages (limitations) of each of the following respirators:

c. Full-face, supplied-air respirator

Atmosphere Supplying Respirators - Supplied Air

Description:

Supplied air respirators use a central source of breathing air that is delivered to the wearer through an air supply line or hose. The respirator type is either a tight-fitting facepiece (half face or full) or loose-fitting hood/suit. There are essentially two major groups of supplied air respirators - the air-line device and the hose mask with or without a blower. Hose masks are not used in power reactors; consequently, further discussion will be limited to demand, pressure demand, and continuous flow air line respirators.

In a demand device, the air enters the facepiece only on "demand" of the wearer, i.e., when the person inhales. During inhalation, there is a negative pressure in the mask, so if there is leakage, contaminated air may enter the mask and be inhaled by the wearer. The pressure demand device has a regulator and valve design such that there is a continuous flow (until a fixed static pressure is attained) of air into the facepiece at all times, regardless of the "demand" of the user. The airflow into the mask creates a positive pressure outward. The continuous-flow air line respirator maintains a constant airflow at all times and does not use a regulator, but uses an airflow control valve or orifice which regulates the flow of air. The continuous-flow device creates a positive pressure in the facepiece. At power reactors, virtually all supplied air operations use the continuous-flow mode.

Limitations:

Since the air line respirator provides no protection if the air supply fails, they shall not be used in IDLH atmospheres or for emergency escape or rescue.

The trailing air supply hose severely limits mobility so it may be unsuitable if frequent movement among separated work stations is required.

The length of hose, number of potential users, and pressure of the supply system can reduce the number of allowable users.
Control of the air quality is essential to avoid introduction of hazardous respiratory agents to the wearers breathing zone.

"Bubble suits" can aspirate air into the suit when the wearer lifts his arms. Consequently, the suit must be tested (for PF) for the exact conditions of use.

**Special Considerations:**

In a situation where the air line respirator is a suit, there shall be a standby rescue person equipped with self contained breathing apparatus and communications equipment whenever supplied-air suits are used.

Requirements for use of respirators in "dangerous" atmospheres is specified in 29 CFR 1910.134(e)(3) as follows:

"(3) Written procedures shall be prepared covering safe use of respirators in dangerous atmospheres that might be encountered in normal operations or in emergencies. Personnel shall be familiar with these procedures and the available respirators.

(i) In areas where the wearer, with failure of the respirator, could be overcome by a toxic or oxygen-deficient atmosphere, at least one additional man shall be present. Communications (visual, voice, or signal line) shall be maintained between both or all individuals present. Planning shall be such that one individual will be unaffected by any likely incident and have the proper rescue equipment to be able to assist the other(s) in case of emergency.

(ii) When self-contained breathing apparatus or hose masks with flowers are used in atmospheres immediately dangerous to life or health, standby men must be present with suitable rescue equipment.

(iii) Persons using air line respirators in atmospheres immediately hazardous to life or health shall be equipped with safety harnesses and safety lines for lifting or removing persons from hazardous atmospheres or other equivalent provisions for the rescue of persons from hazardous atmospheres shall be used. A standby man or men with suitable self-contained breathing apparatus shall be at the nearest fresh air base for emergency rescue."

Manufacturers of airline respirators include instructions specifying a range of air required to produce at least the minimum required flow rates (4 CFM for tight fitting facepiece and 6 CFM for hoods). These specifications are based on hose lengths and the number of
sections connected together. Determining if the proper air flow rate is achieved can be complicated by the use of a breathing air manifold supplying more than one user. The following are recommendations which should be considered.

If all the hose lengths and number of hose fittings are the same, then a manifold with a single regulator and pressure gauge is appropriate for ensuring the proper pressure is used. (Note: If the pressure is within the manufacturer's specifications, then the delivery air flow rate should be at least 4 CFM for tight fitting respirators and 6 CFM for hoods).

For situations where each user has different hose lengths, different number of connection or different air pressure requirements then a separate pressure gauge should be used as follows:

The air flow rate should be measured at the end of the breathing tube (i.e., at the delivery end). This air flow rate should be measured using a calibrated rotameter or equivalent air flow measuring device.

To utilize the Protection Factor (PF) of 2,000 assigned to air supplied hoods, a delivery flow rate of at least 6 CFM but not greater than 15 CFM must be obtained. The individual user's air flow valves should not be altered to maintain a minimum delivery flow rate of 6 CFM as this violates the NIOSH/MSHA approval. If through improper worker training, motivation, or supervision or if operation of the device at the maximum flow rate causes discomfort or unacceptable degradation of hearing or ability to communicate when other workplace factors are considered, the respirator should be operated at lower flow rates and a protection factor of only 1,000 assigned to its use. Taping or otherwise securing the airflow valves in the fully open position does not void the NIOSH/MSHA approval provided the valve is not permanently altered or made so that it would be impossible to increase or decrease the air flow by the user.
2.07.04 Describe the advantages and disadvantages (limitations) of each of the following respirators:

d. Self-contained breathing apparatus (SCBA)

Atmosphere Supplying Respirators - Self-Contained Breathing Apparatus (SCBA)

_Description:

The self-contained breathing apparatus (SCBA) allows the user to carry a respirable breathing supply does not need a stationary air source such as a compressor to provide breathable air. The air supply may last from 3 minutes to 4 hours depending on the nature of the device.

There are two groups of SCBAs - the closed circuit and the open circuit.

Another name for closed circuit SCBAs is "rebreathing" device. The air is rebreathed after the exhaled carbon dioxide has been removed and the oxygen content restored by a compressed oxygen source or an oxygen-generating solid. These devices are designed primarily for 1-4 hours use in toxic atmospheres.

An open circuit SCBA exhausts the exhaled air to the atmosphere instead of recirculating it. A tank of compressed air carried on the back, supplies air via a regulator to the facepiece. Because there is no recirculation of air, the service life of the open circuit SCBA is shorter than the closed circuit system. Two types of open circuit SCBA are available, "demand" or "pressure demand."

In a demand SCBA, air flows into the facepiece only on demand of the wearer, i.e., when the person inhales. During inhalation, there is a negative pressure in the mask so if there is leakage, contaminated air can enter the mask and be breathed by the user. It is important to note that a demand-type SCBA does not provide any higher degree of protection against airborne contaminants than an air-purifying respirator with the same facepiece, but it does provide protection against oxygen deficiency. These types of respirators should not be used for emergency use or for escaping from dangerous environments according to existing guidance (NUREG 0041). The pressure demand open circuit SCBA has a regulator and a valve design which maintains a positive pressure in the facepiece at all times regardless of the "demand" of the user. Because of the high degree of protection provided by the pressure-demand SCBA, this type of unit is recommended for emergency
use, escape and rescue. There also exist combination atmosphere supplying respirators which utilize supplied air and an SCBA.

2.07.04 Describe the advantages and disadvantages (limitations) of each of the following respirators:

   e. Combination atmosphere supplying respirators.

Combination Atmosphere Supplying Respirators

Currently, Mine Safety Appliances (MSA) markets two types of combination supplying respirators: the combination pressure demand breathing apparatus and the dual purpose breathing apparatus. The Combination Pressure Demand Breathing Apparatus marketed by MSA provides respiratory protection for personnel who must work in atmospheres that are immediately dangerous to life or health (IDLH). When connected to a respirable air source, the device permits the wearer to work and move about freely, within the limits of the approved hose length. The Combination Pressure Demand Breathing Apparatus is equipped with a small air cylinder which enables the wearer to escape from dangerous atmospheres in case the primary air supply is interrupted.

The apparatus serves as a long duration work device and as an escape device as well. It is approved for respiratory protection for entry into, for extended periods of work in, and for escape from IDLH atmospheres. If used for entry into IDLH atmospheres, the air line must be connected before entry. The self-contained air supply is approved for escape only.

Operation of the Combination Pressure Demand Breathing Apparatus is manual. It is an approved, rated 5-minute escape device. The pressure demand air line respirator phase is connected by an approved air-supply hose to a primary respirable air source; the worker breathes from this source with the valve of the egress (exit) cylinder of the device turned off until the user is ready to leave the working area. If the primary air supply source should fail for any reason, the worker can switch to the egress cylinder by turning a valve and escape to a safe atmosphere. The worker then can leave, connected to the primary air source, or can open the egress cylinder valve and have approximately five minutes’ respiratory protection. When breathing from the air cylinder, the user can remain connected to the primary air supply and exit, or can disconnect from the air source for easier escape.
The Dual-Purpose Breathing Apparatus combines all the capabilities of a self-contained breathing apparatus and a supplied-air respirator in one unit. The apparatus is approved by the NIOSH and MSHA for use in oxygen deficient atmospheres or where dangerous concentrations of toxic gases or vapors are present. The NIOSH/MSHA approval allows:

- The wearer of the apparatus to enter or exit a dangerous area using only the cylinder air in applications such as emergency rescue
- The wearer to work within the area for a limited time using the cylinder air
- The wearer to work within the area for an extended time using air from a supply line.

Thus, the Dual-Purpose Breathing Apparatus has all the advantages of both Air and Work Masks. Note particularly that 20% of the cylinder air may be used for entry and that the apparatus is not limited to escape. Of course, if the air from the supply line should fail, the wearer can escape the area using the cylinder air.

The Dual-purpose Breathing Apparatus is available in both demand and pressure demand models. In the demand model, air is supplied on demand at ambient atmospheric pressure. In the pressure demand model, a slight positive pressure is maintained within the facepiece during both inhalation and exhalation. The slight positive pressure prevents a toxic atmosphere from leaking into the facepiece; this type of leakage can occur with a demand apparatus due to the negative pressure developed in the facepiece. A pressure demand apparatus should therefore be used where the potential toxicity of the atmosphere is such that no back leakage can be tolerated.

The regulator on the dual purpose breathing apparatus reduces the high pressure from the apparatus's compressed air cylinder to a breathable pressure. In pressure demand models, it also automatically monitors the flow of air into the facepiece so as to maintain a slight positive pressure within the facepiece. The regulator has two inlet ports - one for the cylinder and another for the supply line. A Foster connector allows the air supply to be semi-automatically switched from the cylinder to the air line. With no supply line connected to the regulator, the wearer receives air from the cylinder. When an air line is connected to the regulator through the Foster fitting, the wearer automatically receives air from the supply line. If the air supply from this line should be interrupted, the wearer must disengage the supply line in order to automatically receive air from the cylinder.

**Limitations of the Pressure and Demand SCBA.** The air supply is limited to the amount in the cylinder and therefore the respirator cannot be used for extended periods without recharging or replacing cylinders.
Because these respirators are bulky and heavy, they are often unsuitable for strenuous work or use in confined spaces.

The demand type SCBA work in a negative pressure mode and therefore cannot be used for fire-fighting (10 CFR 50 Appendix R, Section H).

**Special Considerations of the Pressure and Demand SCBA.** As specified in Section 5.5 of NUREG 0041, only the pressure-demand type SCBA should be selected for emergency use, rescue, and re-entry into a contaminated area to perform emergency shutdown or maintenance of equipment.

The performance of SCBAs in high temperature environments, such as fires may lead to rapid deterioration of components.

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### PROTECTION FACTORS

The overall protection afforded by a given respirator design is defined in terms of its protection factor (PF). The PF is defined as *the ratio of the concentration of contaminant in the atmosphere to the concentration inside the facepiece or hood under conditions of use.*

Protection Factors may not be appropriate where chemical or other respiratory hazards exist in addition to radioactive hazards or where the mode of entry is through the skin and not through inhalation. For example, 50% of the intake from exposure to tritiated oxide is through skin absorption. The use of atmosphere supplying respirators will only provide a PF of 2.

Application of PFs is relatively straight forward. The work area airborne radioactivity concentration is divided by the PF to estimate the inhaled concentration. For example, a worker performing steam generator eddy current testing with a full facepiece continuous air flow air line respirator and in an atmosphere of $1 \times 10^6 \mu$Ci/cc Co-60 would be estimated to inhale a concentration of $5 \times 10^{-10} \mu$Ci/cc Co-60. From another perspective, if one wanted to maintain the estimated inhalation concentration at $1 \times 10^{-9} \mu$Ci/cc for an atmosphere of $1 \times 10^{-6} \mu$Ci/cc then the worker must use a respiratory device with a PF of at least 1000.
2.07.06 State the difference between a qualitative and quantitative fit test.

Definitions:

Qualitative fit test: Test to determine if there is any mask leakage, usually using irritant smoke ("Go/no-go" test but no measured value is assigned).

Quantitative fit test: Test to determine quantity of mask leakage and assign a "fit factor," corn oil is the typical challenge atmosphere used (Measures concentration in mask due to leakage against concentration in atmosphere).

2.07.07 State the recommended physical functions the subject must perform during a respirator fit test.

It is impractical to perform a quantitative fit test prior to each entry requiring respiratory protection. Therefore, qualitative tests are performed to ensure an adequate fit for the user. Qualitative tests can use challenge atmospheres such as Isoamyl Acetate (banana oil) or irritant smoke (e.g., stannic chloride) or as a negative or positive pressure test. The irritant smoke test is the most effective since the wearer's obvious discomfort from the smoke will show leakage through the respirator face seal. However, the test produces noxious odors for not only the wearer but those in the test area. The use of "banana oil" requires a subjective evaluation by the wearer and more often than not a user will not admit that in-leakage has occurred. One reactor respiratory program was faithfully utilizing the banana oil to perform the fit test and virtually all wearers indicated no in-leakage through the facepiece. Unfortunately, the respirator only contained a particulate filter cartridge rather than an organic vapor cartridge. Since most reactors use respirators at many different locations, challenge atmosphere tests are difficult to perform and therefore the "immediately prior-to-use" qualitative test normally selected is to perform a negative pressure test.
Additional factors to be considered in fit testing acceptance criteria are the use of communication devices or sorbent canisters with respirators. The respirator approval is voided if any communication device is attached to the facepiece, unless the device is listed in the NIOSH/MSHA approval sheet. MSA "Voice-Con" is one such device that has been approved. Since the combination filter/sorbent, canisters, such as the GMR-II, weigh considerably more than the particulate filter cartridge, the individual should be tested with the canister and not a particulate filter cartridge.

In addition to fit testing personnel, the respirator face pieces and cartridges must be periodically tested. Common practices are to test a portion of particulate cartridges upon procurement and to test all particulate cartridges prior to re-use. Anytime the filter is used by a different individual or on a different day by the same individual, the filter is considered as being reused and should be tested for efficiency, resistance and radioactive contaminants. As long as the exhalation valve for the respirator is in place and functions normally, concern for biological contaminants is of the filter is minimal. Respirator facepieces are tested annually using a test head mannequin and a challenge atmosphere with a light scattering photometer. The challenge atmosphere for testing filters must consist of monodisperse 0.3 micron DPO thermally generated aerosol. The acceptance criteria used is to assume the facepiece seal is almost perfect and any penetration into the breathing zone is from the filter cartridge. Usually a penetration value of less than or equal to 0.003% is considered acceptable.

The subject performs the following functions during fit testing:

1) Normal breathing
2) Deep breathing
3) Moving head from side to side
4) Moving head up and down
5) Frown
6) Talking
7) Running in place
8) Normal breathing
2.07.08 State how the term protection factor (PF) is applied to selection of respiratory protection equipment.

SELECTION

In protecting against radiological airborne contaminants the most critical factor will be meeting the provisions of DOE Order 5480.11 which requires the protection factor for the respirator device used to be greater than the ratio of the work area concentration to the associated DAC.

Additionally, equipment selected must be certified by NIOSH/MSHA or specifically authorized by DOE. Approvals for respiratory devices are authorized in accordance with 30 CFR 11 and the device, type and certification number are listed in NIOSH Publication No. 76-45.

2.07.09 State the general considerations and considerations for the nature of the hazard when selecting the proper respiratory protection equipment.

Selection of the proper respirator for any given situation shall require consideration of the following:

- The nature of the hazard
- The characteristics of the hazardous operation or process
- The location of the hazardous area with respect to a safe area having respirable air
- The period of time for which respiratory protection may be provided
- The activity of the workers in the hazardous area
- The physical characteristics, functional capabilities, and limitations of respirators of various types
- The respirator-protection factors and respirator fit

The following factors concerning the nature of the hazard requiring the use of respirators shall be considered in respirator selection:

- The type of hazard
- Oxygen deficiency
- Contaminant

• The physical and chemical properties
• The physiological effects on the body
• The peak and average concentrations of toxic material or airborne radioactivity level
• The established permissible time-weighted average or peak concentration of toxic material, or both, or established maximum permissible airborne radioactivity level for radioactive substances
• Whether the hazard is an immediately-dangerous-to-life-or-health concentration of toxic material
• Warning properties

Recognition and evaluation of the respiratory hazard (oxygen deficiency or contaminant(s)) shall be an essential part of selecting a respirator except in emergency or rescue operations. Initial monitoring of the respiratory hazard shall be carried out to obtain data needed for the selection of proper respiratory protection. The data should include:

• Identification of the type of respiratory hazard
  - Oxygen deficiency
  - Specific contaminants
• Nature of contaminants
  - Particulate matter
  - Vapors or Gases
• Concentration of respiratory hazard

The following factors concerning the hazardous operation or process shall be taken into account in selecting the proper respirator:

• Operation, process, and work-area characteristics
• Materials, including raw materials, end products, and byproducts (actual and potential)
• Worker activities (Modification in the operation or process shall be taken into account, since this may change the hazard and hence require the selection of a different respirator.)

Selection of air-line respirators includes not only the PF but also the air supply pressure, the air flow to the user and hose length. Each manufacturer's approval sheet lists the approved criteria. For use of 15 to 50 ft of hose at 16 to 20 pounds per square inches, an airflow of greater than 4 CFM to a facepiece, 6 CFM to a hood, and less than 15 CFM to
2.07 - RESPIRATORY PROTECTION

either must be obtained. As discussed, air flow rate delivery should be evaluated for multiple personnel use of breathing air manifolds.

2.07.10 Identify the types of respiratory equipment available for use at your site.

SITE RESPIRATORY EQUIPMENT

(Insert site specific material here)

2.07.11 Identify the quality specifications breathing air must meet.

AIR QUALITY TESTING

An air quality testing program for all sources of respirable air is required. Compressed breathing air shall meet at least the quality specification for Grade D breathing air as described in Compressed Gas Association Commodity Specification G-7.1-1973 (also ANSI Z86.1-1973).

Section 5 of G-7.1 provides acceptable analytical procedures for measuring the respirable air components. Oxygen is easily measured using standard oxygen detectors which utilize an electrolytic reaction to generate a current proportional to the oxygen content. However, a number of reactors perform the measurement incorrectly as the oxygen percentage is determined by the partial pressures of the oxygen in the monitored atmosphere versus the calibrated atmosphere. The test is often performed by placing the detector probe directly in line with the pressurized supply line. Since the air is measured at an increased pressure, the partial pressure will appear greater relative to the calibration partial pressure and an overestimate of the oxygen concentration will result. A better method is to sample the oxygen in a plastic bag and then insert the probe and withdraw the air at a reduced pressure condition. Carbon dioxide and carbon monoxide are easily evaluated using either in line continuous monitors or grab sample "indicator tubes". Draeger sells a portable test kit that consists of indicator tubes which provide a relatively quick assessment of the air quality.
The test for condensed hydrocarbons is usually performed by filtering the air, weighing the filter and calculating the mg/M by assuming the additional filter weight is due to condensed hydrocarbons. For service air systems, the air quality tests should also include monitoring for radioactive contaminants. The test for radioactive contaminants is necessary as a number of service air and breathing air systems have been cross contaminated from radioactive waste or auxiliary boiler contaminants.

The frequency of performing air quality tests is not specified by regulation or in standards. For bottled air systems, such as SCBAs or respirator air supply cylinders, the tests should be performed on a representative sample of the bottles upon receipt at the facility. For facilities which generate respirable air and fill their own SCBA's, the sampling should be performed prior to each lot fill, once during the lot fill and once upon completion of the lot fill. For compressed air supply systems such as fixed station breathing air systems the sampling frequency is best performed prior to each use of a specific manifold system. However, this may be impractical during a major refueling outage where supplied respirable air is extensively used at different stations. In cases of heavy usage, then a daily check of the system may be more appropriate.

**SORBENTS AND PROTECTION AGAINST RADIOIODINES**

The regulations specifically prohibit the use of Pfs for canister sorbents as protection against radioiodine atmospheres. However, the charcoal canisters provided by most manufacturers do provide a measure of protection against radioiodine atmospheres. The efficiency of the charcoal canister is dependent upon the chemical form of the radioiodine, humidity of the atmosphere, and breathing rate of the user. Approval can be obtained from the NRC to use PF's for sorbent cartridges. The criteria for testing and certifying the charcoal cartridges is contained in NUREG/CR-3403, "Criteria and Test Methods for Certifying Air-Purifying Respirator Cartridges and Canisters Against Radioiodine." A brief summary of test conditions and acceptance criteria are as follows in table 1:
Table 1 - Test Conditions and Acceptance Criteria

<table>
<thead>
<tr>
<th>Test Parameter</th>
<th>Criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vapor Concentration</td>
<td>CH$_2$I</td>
</tr>
<tr>
<td>Temperature</td>
<td>1 ppm</td>
</tr>
<tr>
<td>Total Airflow</td>
<td>30 + 1°C</td>
</tr>
<tr>
<td>Equilibration</td>
<td>64 L/min</td>
</tr>
<tr>
<td>(6H at 64 L/min)</td>
<td>All as received</td>
</tr>
<tr>
<td>Maximum Penetration</td>
<td>.01 PPM</td>
</tr>
<tr>
<td>Minimum Service Life</td>
<td>30 min at 100% RH (extrapolated) 60 min at 75% RH</td>
</tr>
</tbody>
</table>

Limiting conditions of use:

Total challenge in the work place (radioactive iodine, non-radioactive iodine or the halogenated compounds) may not exceed 1 ppm.

Temperature in the work area may not exceed 100 °F. Temperature to be measured on each shift or in conjunction with operations which produce heat in the work area.

Respirator wearers must have demonstrated a fit factor greater than 100 on the full facepiece respirator type to which the GMR-1 is attached.

Service life is 8 hours maximum. This is calculated from the time the canister is unsealed and includes periods of non-use. Once the screw cap on the canister threads or the tape seal over the inlet port on the bottom are removed, the 8 hour use duration begins whether used or not by an individual.

Canisters will not be used in the presence of organic solvents, vapors, or chemicals (such as decontamination compounds, lubricants, volatilized paint, alcohol, freon) which could cause aging, poisoning or desorption of the adsorbed radioiodines. Non-exposure to these organic agents must be demonstrated by usage restrictions and by air sampling.

Canisters must be stored in sealed humidity-barrier packaging in a cool, dry environment (QA Class "A" storage).
COMMUNICATIONS

Although conventional respirators distort the human voice to some extent, adequate communication can be maintained in relatively quiet areas. For power reactors, those areas requiring the greatest use of respiratory protection are often the noisiest due to the numerous pumps, motors and fans. Consequently, special attachments or modifications to the respiratory device are often needed to ensure adequate communication.

A mechanical speech-transmission device, called a speaking diaphragm, is an integral part of the facepiece in some respirators. It usually consists of a resonant cavity and diaphragm which transmit sound. The diaphragm also acts as a barrier to the ambient atmosphere and thus should be handled carefully to prevent possible puncture which would permit leakage of an air contaminant into the respirator. Various methods of electronically transmitting and amplifying speech through the respirator are available. These utilize a microphone connected to a speaker, telephone, or radio transmitter. Usually, the microphone is mounted inside the respiratory-inlet covering, while the amplifier, power pack, and speaker or transmitter are attached to the exterior of the respiratory-inlet covering, carried on the body, or remotely located. Respirators with electronic speech-transmission devices having a battery power supply should be used with caution in explosive atmospheres. Sealed power sources shall be checked for integrity of the seals. Connecting cables from the microphones inside the respiratory inlet covering shall have gas-tight seals where they pass through the covering. When the speaker diaphragm is part of the barrier between the respirator wearer and the ambient atmosphere, it shall be and should be adequately protected from puncture or rupture. A microphone mounted on the respirator wearer's throat or head or a microphone/speaker worn in the respirator wearer's ear does not require penetration of a respirator facepiece by a cable.

Any communication device that is an integral part of the respirator or is attached to the exterior such as a sound transducer on the face plate, must be part of the NIOSH/MSHA approval for the respiratory device. One system with a NIOSH/MSHA approval extension for use with negative pressure air purifying respirators is the MSA ClearComV. The ClearComV is a battery operated device with the microphone inserted as an integral part of the facepiece and a plug.
SUMMARY

All respiratory protection devices share a common limitation for protection against hazardous substances which injure the skin or eyes (except SCBAs) or are absorbed through the skin. When selecting any protective device, the chemical form of the hazardous substance should be ascertained to determine if skin protection is required and if the eye protection afforded by the respirator is adequate.

REFERENCES

5. "Limits for Inhalation of Radon Daughters by Workers", ICRP Publication 32.