



BSR/ASHRAE/ASHE Addendum j
to ANSI/ASHRAE/ASHE Standard 170-2017

Public Review Draft

Proposed Addendum j to Standard 170-2017, Ventilation of Health Care Facilities

First Public Review (June 2019)
(Draft shows Proposed Changes to Current Standard)

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FOREWORD

Proposed Addendum j continues the process of reorganizing the standard into three components—Hospital, Outpatient, and Residential Health Care and Support in alignment with the FGI Guidelines’ transition to three separate standards. The intent is not to create additional requirements for outpatient or residential facilities but to separate these from hospital requirements and thus eliminate confusion over which requirements apply to which occupancies. The result will be clarification of a lower level of requirements for outpatient and residential health facilities by separating these from the higher requirement of inpatient facilities. Proposed Addendum j follows the continuing maintenance process in further coordination with FGI staff and 170 staff to result in a coordinated document for use by all stakeholders in the Healthcare Community.

This proposed addendum is the entire Chapter 8 and incorporates Addendum ‘n’. Generally, the changes are as follows:

- *Incorporate Addendum ‘a’ updated filtration requirements.*
- *Revise the space name terminology, table organization, and subheadings to better correlate with the 2018 FGI Guidelines for Design and Construction of Hospitals and Outpatient Facilities, including the addition of paragraph numbers after each space name. These revised 2018 FGI paragraph numbers have been coordinated with FGI committee members and will be presented in italicized text to keep them as informative language.*
- *Several spaces have relocated within the two Tables (Table 8.1 & Table 8.2). Table 8.1 and notes are reorganized into a new Table 8.1 and notes and new Table 8.2 and notes as shown; cyan highlighted spaces move to new Table 8.1 and notes and green highlighted spaces move to new Table 8.2 and notes, otherwise Table changes are shown in underline (new Table) and strikethrough (old Table).*

Note: In this addendum, generally, changes to the current standard are indicated in the text by underlining (for additions) and ~~strikethrough~~ (for deletions) unless the instructions specifically mention some other means of indicating the changes.

Addendum j to 170-2017

Revise Section 8 as shown. Only changes are shown in underline and strikethrough.

8. SPACE VENTILATION—OUTPATIENT SPACES

The ventilation requirements of this standard are minimums that provide control of environmental comfort, asepsis, and odor in outpatient spaces. However, because they are minimum requirements and because of the diversity of the population and variations in susceptibility and sensitivity, these requirements do not provide assured protection from discomfort, airborne transmission of contagions, and odors.

8.1 General Requirements. Specialized Outpatient Facility Requirements. The following facility types shall comply with this section; Outpatient Surgical, Endoscopy, Infusion, Renal Dialysis, Freestanding Emergency Departments and Imaging Facilities with Class 2 and 3 Imaging Rooms. The following general requirements shall apply for space ventilation:

- a. Spaces shall be ventilated according to Table 8.1.
 1. Design of the ventilation system shall provide air movement that is generally from clean to less-clean areas. If

any form of variable-air-volume or load-shedding system is used for energy conservation, it shall not compromise the pressure balancing relationships or the minimum air changes required by the table.

2. The ventilation requirements in this table are intended to provide for comfort as well as for asepsis and odor control in spaces of a health care facility that directly affect patient care. For spaces not specifically listed here, ventilation requirements shall be that of functionally equivalent spaces in the table. If no functionally equivalent spaces exist in the table, ventilation requirements shall be obtained from ANSI/ASHRAE Standard 62.1¹. Where spaces with prescribed rates in both Standard 62.1 and Table 8.1 of this standard exist, the higher of the two air change rates shall be used.
3. For design purposes, the minimum number of total air changes indicated shall be either supplied for positive pressure rooms or exhausted for negative pressure rooms. Spaces that are required in Table 8.1 to be at a negative pressure relationship and that are not required to be exhausted shall use the supply airflow rate to compute the minimum total air changes per hour required. For spaces that require a positive or negative pressure relationship, the number of air changes can be reduced when the space is unoccupied, provided that the required pressure relationship to adjoining spaces is maintained while the space is unoccupied and that the minimum number of air changes indicated is reestablished anytime the space becomes occupied. Controls intended to switch the required pressure relationships between spaces from positive to negative, and vice versa, shall not be permitted. Air change rates in excess of the minimum values are expected in some cases in order to maintain room temperature and humidity conditions based on the space cooling or heating load.
4. The entire minimum outdoor air changes per hour required by Table 8.1 for the space shall meet the filtration requirements of Section 8.1.
5. For spaces where Tables 8.1 permits air to be recirculated by room units, the portion of the minimum total air changes per hour required for a space that is greater than the minimum outdoor air changes per hour required component may be provided by recirculating room HVAC units. Such recirculating room HVAC units shall
 - i. not receive nonfiltered, nonconditioned outdoor air;
 - ii. serve only a single space; and
 - iii. provide a minimum MERV 6 filter for airflow passing over any surface that is designed to condense water. This filter shall be located upstream of any such cold surface, so that all of the air passing over the cold surface is filtered.
6. For air-handling systems serving multiple spaces, system minimum outdoor air quantity shall be calculated using one of the following methods:
 - i. System minimum outdoor air quantity for an air-handling system shall be calculated as the sum of the individual space requirements as defined by this standard.
 - ii. System minimum outdoor air quantity shall be calculated by the Ventilation Rate Procedure (multiple zone formula) of ASHRAE Standard 62.1¹. The minimum outdoor air change rate listed in this standard shall be interpreted as the V_{Oz} (zone outdoor airflow) for purposes of this calculation.
- b. Air filtration for spaces shall comply with Section 6.4 and Tables 8.1.
- c. Supply air outlets for spaces shall comply with Table 6.7.2.
- d. In All rooms, protective environment rooms, operating and procedure rooms, heating with supply air or radiant panels that meet the requirements of Section 6.5.3 shall be provided.
- e. In a building that contains a mixture of spaces programmed for outpatient care as well as spaces programmed for inpatient care, the outpatient care spaces shall be designed in accordance with Table 8.1, and the inpatient care spaces shall be designed in accordance with Table 7.1.

Table 8.1 and notes are reorganized into a new Table 8.1 and notes and new Table 8.2 and notes as shown; cyan highlighted spaces move to new Table 8.1 and notes and green highlighted spaces move to new Table 8.2 and notes, otherwise Table changes are shown in underline (new Table) and strikethrough (old Table).

Table 8.1 Design Parameters for Outpatient Specific Spaces

Function of Space (f)	Pressure Relationship to Adjacent Areas (n)	Minimum Outdoor ach	Minimum Total ach	All Room Air Exhausted Directly to Outdoors (j)	Air Recirculated by Means of Room Units (a)	Minimum Filter Efficiencies (c)	Design Relative Humidity (k), %	Design Temperature (l), °F/°C
COMMON SPACES IN OUTPATIENT FACILITIES								
All anteroom (i) (3.1-3.4.3)	(e)	NR	10	Yes	No	7/NR	NR	NR
All room (i) (3.1-3.4.2)	Negative	2	12	Yes	No	7/NR	Max 60	70-75/21-24
Bronchoscopy, sputum collection, and pentamidine administration (n)	Negative	2	12	Yes	No	7/NR	NR	68-73/20-23
Clean supply storage (3.1-3.6.9)	Positive	2	4	NR	NR	7/NR	Max 60	72-78/22-26
Emergency waiting rooms	Negative	2	12	Yes (q)	NR	7/NR	Max. 65	70-75/21-24
Environmental services room (3.1-5.5.1)	Negative	NR	10	Yes	No	7/NR	NR	NR
General purpose examination/observation room (3.1-3.2.2)	NR	2	4	NR	NR	7/NR	Max 60	70-75/21-24
Laboratory testing/work area if in a separate dedicated room (3.1-4.1.2)	Negative	2	6	Yes	NR	7/NR	NR	70-75/21-24
Medical waste holding spaces (3.1-5.4.1.3)	Negative	2	10	Yes	No	7/NR	NR	NR
Medication preparation room programmed to compound sterile preparations (b) (3.1-3.6.6.2)	Positive	2	4	NR	NR	7/HEPA (s)	NR	NR
Soiled holding room (3.1-3.6.10)	Negative	2	6	Yes	No	7/NR	NR	72-78/22-26
Special purpose examination room (3.1-3.2.3)	NR	2	6	NR	NR	7/NR	Max 60	70-75/21-24
SPACES SPECIFIC TO PARTICULAR OUTPATIENT FACILITIES								
Cancer treatment area (p) (3.6-3.2)	NR	2	6	NR	NR	7/NR	Max 60	70-75/21-24
Diagnostic imaging waiting area (3.5-6.1.3.2) (g)	Negative	2	12	Yes (q), (r)	NR	7/NR	Max 60	70-75/21-24
ECT procedure room (p) (3.11-3.3.2.2)	NR	2	4	NR	NR	7/NR	Max 60	70-75/21-24
Endoscopy procedure room (h) (3.9-3.2.2)	NR	2	6	NR	No	7/NR	Max 60	68-73/20-23
Freestanding urgent care facility procedure room (3.5-3.2.2)	Positive	2	6	NR	No	7/NR	NR	70-75/21-24
Instrument processing room (3.9-5.1)	Negative	2	10	Yes	No	7/NR	NR	NR
Office based procedure room (p) (3.8-3.1)	NR	2	4	NR	NR	7/NR	Max 60	70-75/21-24
Outpatient surgical facility operating room (m), (o) (3.7-3.3)	Positive	4	20	NR	No	7/14	20-60	68-75/20-24
Outpatient surgical facility procedure room (o), (d) (3.7-3.2)	Positive	3	15	NR	No	7/NR	20-60	70-75/21-24
Postoperative recovery area (3.7-3.4.3)	NR	2	6	NR	No	7/NR	Max 60	70-75/21-24
Postprocedure recovery area (u) (3.9-3.3)	NR	2	2	NR	NR	7/NR	Max 60	70-75/21-24
Preprocedure patient care area (t) (3.9-3.3)	NR	2	2	NR	NR	7/NR	Max 60	70-75/21-24

Note: NR = no requirement

Table 8.1 Design Parameters—Specialized Outpatient Spaces

Function of Space (f)	Pressure Relationship to Adjacent Areas (n)	Minimum Outdoor ach	Minimum Total ach	All Room Air Exhausted Directly to Outdoors (j)	Air Recirculated by Means of Room Units (a)	Minimum Filter Efficiencies (c)	Design Relative Humidity (k), %	Design Temperature (l), °F/°C
<u>SURGERY AND EMERGENCY DEPT (ED)</u>								
<u>Delivery (Caesarean) (m), (o), (v) (FGI 2.1-3.2.3)</u>	Positive	4	20	NR	No	MERV-A-16 (dd)	20–60	68–75/20–24
<u>ED human decontamination (FGI 2.8-3.4.8)</u>	Negative	2	12	Yes	No	MERV-A-14 (cc)	NR	NR
<u>ED exam/treatment room (p) (FGI 2.8-3.4.2)</u>	NR	2	6	NR	NR	MERV-A-14 (cc)	Max 60	70–75/21–24
<u>ED public waiting area (FGI 2.8-6.2.3)</u>	Negative	2	12	Yes (q)	NR	MERV-A-8	Max 65	70–75/21–24
<u>Operating room (m), (o), (v) (FGI 2.1-3.2.3)</u>	Positive	4	20	NR	No	MERV-A-16 (dd)	20–60	68–75/20–24
<u>Procedure room (d), (p) (FGI 2.1-3.2.2)</u>	Positive	3	15	NR	No	MERV-A-14	20–60	70–75/21–24
<u>Phase I recovery (PACU) (FGI 2.1-3.7.4)</u>	NR	2	6	NR	No	MERV-A-8	Max 60	70–75/21–24
<u>Phase II recovery (u) (FGI 2.1-3.7.5)</u>	NR	2	2	NR	NR	MERV-A-8	Max 60	70–75/21–24
<u>Pre-procedure patient care (t) (FGI 2.1-3.7.3)</u>	NR	2	2	NR	NR	MERV-A-8	Max 60	70–75/21–24
<u>Trauma room (crisis or shock) (bb) (FGI 2.8-3.4.4)</u>	Positive	3	15	NR	No	MERV-A-14	20–60	70–75/21–24
<u>Triage (FGI 2.8-6.2.2.2 & 6.2.2.3)</u>	Negative	2	12	Yes (q)	NR	MERV-A-8	Max 60	70–75/21–24
<u>DIAGNOSTIC AND TREATMENT</u>								
<u>Class 1 imaging room (FGI 2.1-3.5.2.4(1)(b)(i))</u>	NR	2	6	NR	NR	MERV-A-8	Max 60	70–75/21–24
<u>Class 2 imaging room (d), (p) (FGI 2.1-3.5.2.4(1)(b)(ii))</u>	Positive	3	15	NR	No	MERV-A-14	20–60	70–75/21–24
<u>Class 3 imaging room (m), (o) (FGI 2.1-3.5.2.4(1)(b)(iii))</u>	Positive	4	20	NR	No	MERV-A-16 (dd)	20–60	68–75/20–24
<u>Diagnostic imaging waiting (g) (FGI 2.1-3.5.10.4)</u>	Negative	2	12	Yes (q), (r)	NR	MERV-A-8	Max 60	70–75/21–24
<u>All anteroom (i) (FGI 2.1-3.3.2.3)</u>	(e)	NR	10	Yes	No	MERV-A-8	NR	NR
<u>All room (i) (FGI 2.1-3.3.2)</u>	Negative	2	12	Yes	No	MERV-A-8	Max 60	70–75/21–24
<u>PE anteroom (n) (w) (FGI 1.2-4.2.2.1(1))</u>	(e)	NR	10	NR	No	HEPA	NR	NR
<u>Protective environment room (n) (w) (FGI 1.2-4.2.2.1(1))</u>	Positive	2	12	NR	No	HEPA	Max 60	70–75/21–24
<u>Cancer treatment area (FGI 2.6-3.1)</u>	NR	2	6	NR	NR	MERV-A-8	Max 60	70–75/21–24
<u>Dialysis treatment area (FGI 2.10-3.2)</u>	NR	2	6	NR	NR	MERV-A-8	NR	72–78/22–26
<u>Dialyzer reprocessing room (FGI 2.10-3.8.12)</u>	Negative	NR	10	Yes	No	MERV-A-8	NR	NR
<u>Bronchoscopy (n) (x) (FGI 2.1-3.2.2.1)</u>	Negative	2	12	Yes	No	MERV-A-14	NR	68–73/20–23
<u>Instrument processing room (FGI 2.1-4.3.2.3)</u>	Negative	2	10	Yes	No	MERV-A-8	NR	NR
<u>Endoscopy procedure room (h) (FGI 2.9-3.2)</u>	NR	2	6	NR	No	MERV-A-8	Max 60	68–73/20–23

Note: NR = no requirement

Table 8.1 Design Parameters—Specialized Outpatient Spaces (Continued)

Function of Space (f)	Pressure Relationship to Adjacent Areas (n)	Minimum Outdoor ach	Minimum Total ach	All Room Air Exhausted Directly to Outdoors (j)	Air Recirculated by Means of Room Units (a)	Minimum Filter Efficiencies (c)	Design Relative Humidity (k), %	Design Temperature (l), °F/°C
<u>DIAGNOSTIC AND TREATMENT (Continued)</u>								
Examination/observation (<i>FGI 2.1-3.2.1</i>)	NR	2	4	NR	NR	<u>MERV-A-8</u>	Max 60	70–75/21–24
Specialty IC exam room (y) (<i>FGI 2.1-3.2.1.3</i>)	<u>Negative</u>	2	6	<u>Yes</u>	NR	<u>MERV-A-8</u>	Max 60	70–75/21–24
Laboratory work room (z) (<i>FGI 2.1-4.1.2.1</i>)	<u>Negative</u>	2	6	<u>Yes</u>	NR	<u>MERV-A-8</u>	NR	70–75/21–24
Pharmacy/Med Prep (b) (<i>FGI 2.1-3.8.8.2 & 2.1-4.2.2</i>)	Positive	2	4	NR	NR	<u>MERV-A-8</u>	NR	NR
Laser eye room (<i>FGI 2.1-3.2.2</i>)	<u>NR</u>	<u>2</u>	<u>6</u>	<u>NR</u>	<u>No</u>	<u>MERV-A-8</u>	<u>Max 60</u>	<u>68–73/20–23</u>
Nuclear medicine (see Section 8.7) (<i>FGI 2.1-3.5.7</i>)	<u>Negative</u>	<u>2</u>	<u>6</u>	<u>Yes</u>	<u>No</u>	<u>MERV-A-8</u>	<u>NR</u>	<u>70–75/21–24</u>
Toilet or Toilet/Shower room (<i>FGI 2.1-3.10.2</i>)	<u>Negative</u>	<u>NR</u>	<u>10</u>	<u>Yes</u>	<u>No</u>	<u>MERV-A-8</u>	<u>NR</u>	<u>NR</u>
<u>STERILE PROCESSING (aa)</u>								
One-room sterile processing (<i>FGI 2.1-4.3.2.3</i>)	<u>NR</u>	<u>2</u>	<u>6</u>	<u>NR</u>	<u>No</u>	<u>MERV-A-14 (ee)</u>	<u>NR</u>	<u>NR</u>
Sterilizer equipment room (<i>FGI 2.1-4.3.2.2</i>)	<u>Negative</u>	<u>NR</u>	<u>10</u>	<u>Yes</u>	<u>No</u>	<u>MERV-A-8</u>	<u>NR</u>	<u>NR</u>
Clean workroom (<i>FGI 2.1-4.3.2.2.3</i>)	<u>Positive</u>	<u>2</u>	<u>4</u>	<u>NR</u>	<u>No</u>	<u>MERV-A-14 (ee)</u>	<u>Max 60</u>	<u>60–73/16–23</u>
Clean supply storage (<i>FGI 2.1-4.3.2.2.4</i>)	Positive	2	4	NR	NR	<u>MERV-A-14 (ee)</u>	Max 60	72–78/22–26
Supply Receiving (<i>FGI 2.1-4.3.2.4</i>)	<u>Negative</u>	<u>NR</u>	<u>10</u>	<u>Yes</u>	<u>No</u>	<u>MERV-A-8</u>	<u>NR</u>	<u>NR</u>
Decontamination room (<i>FGI 2.1-4.3.2.2</i>)	<u>Negative</u>	<u>2</u>	<u>6</u>	<u>Yes</u>	<u>No</u>	<u>MERV-A-8</u>	<u>NR</u>	<u>60–73/16–23</u>
<u>SERVICE / SUPPORT SPACE</u>								
Environmental services room (<i>FGI 2.1-5.3.1</i>)	<u>Negative</u>	NR	10	<u>Yes</u>	No	<u>MERV-A-8</u>	NR	NR
Laundry/Linen Processing (<i>FGI 2.1-4.4.2.1</i>)	<u>Negative</u>	<u>2</u>	<u>10</u>	<u>Yes</u>	<u>No</u>	<u>MERV-A-8</u>	<u>NR</u>	<u>NR</u>
Clean workroom or clean supply (<i>FGI 2.1-3.8.11</i>)	<u>Positive</u>	<u>2</u>	<u>4</u>	<u>NR</u>	<u>NR</u>	<u>MERV-A-8</u>	<u>NR</u>	<u>NR</u>
Regulated waste holding (<i>FGI 2.1-5.2.1.3</i>)	<u>Negative</u>	2	10	<u>Yes</u>	No	<u>MERV-A-8</u>	NR	NR
Soiled workroom or soiled holding (<i>FGI 2.1-3.8.12</i>)	<u>Negative</u>	2	6	<u>Yes</u>	No	<u>MERV-A-8</u>	NR	72–78/22–26

Note: NR = no requirement

Normative Notes for Table 8.1:

- a. Except where indicated by a “No” in this column, recirculating room HVAC units (with heating or cooling coils) are acceptable for providing that portion of the minimum total air changes per hour that is permitted by Section 8.1 (subparagraph [a][5]). Because of the cleaning difficulty and potential for buildup of contamination, recirculating room units shall not be used in areas marked “No.” Recirculating devices with high-efficiency particulate air (HEPA) filters shall be permitted in existing facilities as interim, supplemental environmental controls to meet requirements for the control of airborne infectious agents. The design of either portable or fixed systems should prevent stagnation and short circuiting of airflow. The design of such systems shall also allow for easy access for scheduled preventative maintenance and cleaning.
- b. Pharmacy compounding areas may have additional air change, differential pressure, and filtering requirements beyond the minimum of this table, depending on the type of pharmacy, the regulatory requirements (which may include adoption of USP 795-General Pharmacy/Nonsterile Preparations, USP 797-Sterile Compounding and USP 800-Hazardous Drugs *toxins*), the associated level of risk of the work, and the equipment used in the spaces. ***Informative Note:*** See USP (2012) in Appendix B. ***Informative Notes:*** 1) See USP (2017a & b) in Appendix B. 2) Hazardous Drug Sterile Compounding follows USP-797 and then applies USP-800 as an overlay of requirements (e.g.-Hazardous Drug storage follows USP-800 only).
- c. Table entries are the minimum filter efficiencies required for the space. Refer to Section 6.4 of this document for further clarification of filtration requirements. ~~The first table entry is the minimum filter efficiency for Filter Bank No. 1. The second table entry (after the slash) is the minimum filter efficiency for Filter Bank No. 2. The minimum efficiency reporting value (MERV) is based on the method of testing described in ANSI/ASHRAE Standard 52.2.~~ ***Informative Note:*** See ASHRAE [2012] in Appendix B. Minimum filter requirements shall meet the equivalent MERV-A rating when tested in accordance with Appendix J of ANSI/ASHRAE Standard 52.2.
- d. Pressure relationships need not be maintained when the room is unoccupied.
- e. See Section 8.2 7.2.1 and its subsections for ventilation requirements including pressure relationship requirements.
- f. Parenthetical notations following a space name are paragraph references to the 20142018 Facility Guidelines Institute document *Guidelines for Design and Construction of Hospitals and Outpatient Facilities (Informative Note: FGI [20142018])*. These FGI paragraph references are provided to the user of the standard to aid in the application of design requirements.
- g. These ventilation requirements only apply to ~~urgent care facility~~ waiting areas where the ICRA determines that the diagnostic imaging waiting area requires special consideration to reduce the risk of airborne infection transmission. If the ICRA does not have these special consideration provisions then the ventilation requirements shall meet the provisions of ANSI/ASHRAE Standard 62.1¹.
- h. If the planned space is designated in the organization’s operational plan to be used for both bronchoscopy and gastrointestinal endoscopy, the design parameters for “bronchoscopy, sputum collection, and pentamidine administration” shall be used.
- i. The AII room described in this standard shall be used for isolating the airborne spread of infectious diseases, such as measles, varicella, or tuberculosis. Supplemental recirculating devices using HEPA filters shall be permitted in the AII room to increase the equivalent room air exchanges; however, the minimum outdoor air changes of Table 8.1 are still required. When the AII room is not used for airborne infection isolation, the pressure relationship to adjacent areas, when measured with the door closed, shall remain unchanged, and the minimum total air change rate shall be 6ach.
- j. In some areas with potential contamination and/or odor problems, exhaust air shall be discharged directly to the outdoors and not recirculated to other areas. Individual circumstances may require special consideration for air exhausted to the outdoors. To satisfy exhaust needs, constant replacement air from the outdoors is necessary when the system is in operation.
- k. The RH ranges listed are the minimum and/or maximum allowable at any point within the design temperature range required for that space.
- l. Systems shall be capable of maintaining the rooms within the range during normal operation. Lower or higher temperature shall be permitted when patients’ comfort and/or medical conditions require those conditions.
- m. National Institute for Occupational Safety and Health (NIOSH) criteria documents⁹ regarding occupational exposure to waste anesthetic gases and vapors and control of occupational exposure to nitrous oxide indicate a need for both local exhaust (scavenging) systems and general ventilation of the areas in which the respective gases are used. Refer to NFPA 99¹⁰ for other requirements.
- n. If pressure-monitoring device alarms are installed, allowances shall be made to prevent nuisance alarms. Short-term excursions from required pressure relationships shall be allowed while doors are moving or temporarily open. Simple visual methods such as smoke trail, ball-in-tube, or flutterstrip shall be permitted for verification of air-flow direction.
- o. Surgeons or surgical procedures may require room temperatures, ventilation rates, humidity ranges, and/or air distribution methods that exceed the minimum indicated ranges.
- p. Treatment or procedure rooms used for bronchoscopy shall be treated as bronchoscopy rooms. Treatment or procedure rooms used for procedures with nitrous oxide shall contain provisions for exhausting anesthetic waste gases.
- q. In a recirculating ventilation system, HEPA filters shall be permitted instead of exhausting the air from these spaces to the outdoors, provided that the return air passes through the HEPA filters before it is introduced into any other spaces. The entire minimum total air changes per hour of recirculating airflow shall pass through HEPA filters. When these areas are open to larger, nonwaiting spaces, the exhaust air volume shall be calculated based on the seating area of the waiting area. ***Informative Note:*** The intent here is to not require the volume calculation to include a very large space (e.g., an atrium) just because a waiting area opens onto it.
- r. The requirement that all room air be exhausted directly to outdoors applies only to radiology waiting rooms programmed to hold patients who are waiting for chest x-rays for diagnosis of respiratory disease.
- s. As an alternative to the requirement for HEPA filters in Filter Bank No. 2, MERV-14 rated filters may be used in Filter Bank No. 2 if a tertiary terminal HEPA filter is provided for this space. ***Informative Note:*** High-efficiency particulate air (HEPA) filters are those filters that remove at least 99.97% of 0.3 micron-sized particles at the rated flow in accordance with the testing methods of IEST RP- CC001.6 See IEST [20052016] in Appendix B.
- t. If anesthetic gases are administered in the area, the minimum total air changes shall be increased to 6.
- u. If anesthetic gases are used during the preceding procedure, the minimum total air changes shall be increased to 6.
- v. See section 7.4.1 of this standard for ventilation requirements. The Operating Room designation includes surgical cystoscopic rooms.

- w. See section 7.2.2 of this standard for ventilation requirements.
- x. This space includes sputum collection and pentamidine administration. See section 8.5.2 of this standard.
- y. Examination rooms (*Identified as Specialty Infection Control-IC Exam rooms*) programmed for use by patients with undiagnosed gastrointestinal symptoms, undiagnosed respiratory symptoms, or undiagnosed skin symptoms.
- z. This room is intended for basic blood & urine specimen testing and short term storage only. Outpatient facilities which provide the following specialized service spaces/rooms should consult Table 7.1 of this standard for ventilation requirements.
 - i. Laboratory work areas including the specialty lab services such as bacteriology, biochemistry, cytology, glasswashing, histology, media transfer, microbiology, nuclear medicine, pathology, serology and/or sterilizing.
- aa. See AAMI Standard ST79¹¹ for additional information for these spaces.
- bb. The term *trauma room* as used herein is a first-aid room and/or emergency department room used for general initial treatment of accident victims. The OR within the trauma center that is routinely used for emergency surgery is considered to be an OR by this standard.
- cc. A minimum MERV-A-8 filter may be utilized for this space in lieu of a minimum MERV-A-14 filter if all room air is exhausted directly to the outdoors and the pressure relationship to adjacent areas is kept negative. If a filter rated less than MERV-A-14 is utilized the space shall be considered “Negative” with regards to the table and must comply with all other requirements for negative spaces within the standard.
- dd. See section 7.4.1.c.
- ee. Minimum MERV-A-14 filters shall be required for spaces where sterile equipment is packed into sterile packages. MERV-A-8 filters may be used in place of MERV-A-14 in spaces where sterile products are stored in sealed packaging but are not opened or otherwise handled outside of the sealed package.

8.2 General Outpatient Facility Requirements. All outpatient facility types other than those indicated in Section 8.1 shall comply with this Section and Table 8.2.

Unless otherwise noted in this section, all requirements for space ventilation of general outpatient spaces are contained within this Section and Table 8.2 and Sections 6, 7, 9 and 10 of this Standard shall not apply. For requirements related to Sections 6 and 10 which are not found in this Section, refer to local and state building codes. Where no local or state code is recognized, the requirements of ANSI/ASHRAE Standard 62.1¹ shall apply.

Informative Note: ASHRAE recognizes that ASHRAE Standards are typically the foundation of state and local building codes. However, state and local codes also represent important regional interests and conditions. As such, state and local building codes shall also be followed to the maximum extent practicable.

The following requirements shall apply for space ventilation:

a. Spaces shall be ventilated according to Table 8.2.

1. Design of the ventilation system shall provide air movement that is generally from clean to less-clean areas. If any form of variable-air-volume or load-shedding system is used for energy conservation, it shall not compromise the pressure balancing relationships or the minimum air changes required by the table.
2. The ventilation rates in this table are intended to provide for comfort as well as for asepsis and odor control in spaces of a health care facility that directly affect patient care. Ventilation rates for clinical spaces not specified here shall be that of functionally equivalent spaces in the table. If no functionally equivalent spaces exist in the table, ventilation requirements shall be obtained from ANSI/ASHRAE Standard 62.1¹. Where spaces with prescribed rates in both Standard 62.1 and Table 8.2 of this standard exist, the higher of the two air change rates shall be used.
3. For design purposes pressure relationships shall be achieved by the following methods:
 - i. Spaces that require a positive or negative pressure relationship shall maintain the required pressure relationship during room occupied hours.
 - ii. For systems utilizing air changes per hour, the minimum number of total air changes indicated shall be either supplied for positive pressure rooms or exhausted for negative pressure rooms. Spaces that are required in Table 8.2 to be at a negative pressure relationship and that are not required to be exhausted shall use the supply airflow rate to compute the minimum total air changes per hour required. For spaces that require a positive or negative pressure relationship, the number of air changes can be reduced when the space is unoccupied, provided that the required pressure relationship to adjoining spaces is maintained while the space is unoccupied and that the minimum number of air changes indicated is reestablished anytime the space becomes occupied. Controls intended to switch the required pressure relationships between spaces from positive to negative, and vice versa, shall not be permitted. Air change rates in excess of the minimum values are expected in some cases in order to maintain room temperature and humidity conditions based on the space cooling or heating load.
4. All ventilation required by Table 8.2 shall meet the filtration requirements of Section 6.4 and Section 8.2.
5. For spaces where Tables 8.2 permits air to be recirculated by room units, the portion of the minimum total air changes per hour required for a space that is greater than the minimum outdoor air changes per hour required component may be provided by recirculating room HVAC units. Such recirculating room HVAC units shall
 - i. not receive nonfiltered, nonconditioned outdoor air;
 - ii. provide the manufacturer's recommended filter (or MERV-A-8 as a minimum) for airflow passing over any surface that is designed to condense water. This filter shall be located upstream of any such cold surface, so that all of the air passing over the cold surface is filtered.
6. For air-handling systems utilizing the CFM/person and CFM/sq. ft. outside air ventilation rates, system minimum outdoor air quantity shall be calculated by the Ventilation Rate Procedure of ASHRAE Standard 62.1¹. The CFM/person rate shall be considered the R_p value and the CFM/sq. ft. rate shall be considered the R_a value in the calculation.
 - i. The minimum occupant quantity is provided as a required minimum in the R_p column of Table 8.2. The design zone population (P_z) shall equal the largest (peak) number of people expected to occupy the ventilation zone during typical use. When the design zone population is less than the minimum occupant quantity provided in Table 8.2, then the occupant factor in the R_p column of Table 8.2 shall be utilized in lieu of the design zone population.
 - ii. A zone minimum primary airflow shall be provided as follows; For each zone, the minimum primary airflow (V_{pz-min}) shall be determined by the following Equation
$$V_{pz-min} = V_{OZ} \times 1.5$$
7. For air-handling systems serving multiple spaces and utilizing the minimum outdoor ach column, system minimum outdoor air quantity shall be calculated using one of the following methods:
 - i. System minimum outdoor air quantity for an air-handling system shall be calculated as the sum of the individual space requirements as defined by this standard.
 - ii. System minimum outdoor air quantity shall be calculated by the Ventilation Rate Procedure (multiple zone formula) of ASHRAE Standard 62.1¹. The minimum outdoor air change rate listed in this standard shall be interpreted as the V_{OZ} (zone outdoor airflow) for purposes of this calculation.
8. Specific only to Table 8.2 design paths. In lieu of calculating ventilation via air change rate (ach), an optional path allows calculating ventilation rate using provided values for

cfm/person (Rp) and cfm/sq.ft. area (Ra) and Air Class. This alternative design path option excludes compliance with the space total air change rate (ach) requirement. This alternative design path option is **only** applicable to Table 8.2 spaces. Note: there is no correlation expected or intended between the two design path options.

Add Table 8.2 and notes as shown.

Table 8.2 Design Parameters—General Outpatient Spaces (iq)

Function of Space (if)	Pressure Relationship to Adjacent Areas (id)	ach design option		All Room Air Exhausted Directly to Outdoors (ij)	AirRecirc-ulated by Means of Room Units (ia)	Minimum Filter Efficiencies (ic)	Design Relative Humidity % (ii)	Design Temperature °F/°C (ik)	Rp-Ra-air class design option		
		Min. Outdoor ach (iq)	Min. Total ach (iq)						Air Class (iq)	Rp Cfm/(Ls)/ Person and Occupant Factor (iq)	Ra Cfm/ft/(Ls/m) (iq)
GENERAL DIAGNOSTIC AND TREATMENT											
Birthing room (FGI 2.4-2.2)	NR	2	3	Yes (ih)	NR	MERV-A-14	Max 60	70–75/21–24	2	10 (5) / 4	0.18 / (0.9)
Urgent care exam (ie) (FGI 2.5-3.2.1)	NR	2	3	NR	NR	MERV-A-8	NR	70–75/21–24	2	7.5 (3.8) / 3	0.12 / (0.6)
Urgent care treatment (ie) (FGI 2.5-3.2.2)	NR	2	3	NR	NR	MERV-A-8	NR	70–75/21–24	2	7.5 (3.8) / 3	0.18 / (0.9)
Urgent care triage (FGI 2.5-3.2.3)	Negative	2	3	Yes	NR	MERV-A-8	Max 60	70–75/21–24	3	10 (5) / 3	0.18 / (0.9)
Urgent care observation (FGI 2.5-3.3)	NR	2	2	NR	NR	MERV-A-8	NR	70–75/21–24	2	5 (2.5) / 2	0.12 / (0.6)
General examination room (FGI 2.1-3.2.1)	NR	2	2	NR	NR	MERV-A-8	NR	70–75/21–24	1	7.5 (3.8) / 3	0.12 / (0.6)
Specialty IC exam room (ib) (FGI 2.5-3.2.3)	Negative	2	3	Yes	NR	MERV-A-8	Max 60	70–75/21–24	3	10 (5) / 3	0.18 / (0.9)
Laboratory work room (il) (FGI 2.1-4.1.2.1)	NR	2	3	Yes (ih)	NR	MERV-A-8	NR	70–75/21–24	2	7.5 (3.8) / 2	0.12 / (0.6)
Medication room (FGI 2.1-3.8.8.2)	Positive	2	2	NR	NR	MERV-A-8	Max 60	70–75/21–24	1	5 (2.5) / 2	0.18 / (0.9)
Class 1 Imaging rooms (ig) (FGI 2.1-3.5)	NR	2	3	NR	NR	MERV-A-8	Max 60	72–78/22–26	1	7.5 (3.8) / 2	0.12 / (0.6)
Psychiatric Examination Room (FGI 2.11-3.2.2)	NR	2	3	NR	NR	MERV-A-8	NR	70–75/21–24	1	5 (2.5) / 2	0.06 / (0.3)
Psychiatric Consultation Room (FGI 2.11-3.2.4)	NR	2	3	NR	NR	MERV-A-8	NR	70–75/21–24	1	5 (2.5) / 2	0.06 / (0.3)
Psychiatric Group Room (FGI 2.11-3.2.5)	NR	2	3	NR	NR	MERV-A-8	NR	70–75/21–24	1	5 (2.5) / 2	0.06 / (0.3)
Psychiatric Seclusion Room (FGI 2.11-3.2.7)	NR	2	2	NR	NR	MERV-A-8	NR	70–75/21–24	2	10 (5) / 3	0.12 / (0.6)
ECT procedure room (FGI 2.11-3.2.9.2)	NR	2	2	NR	NR	MERV-A-8	NR	70–75/21–24	1	7.5 (3.8) / 3	0.12 / (0.6)
Physical Therapy Individual Room (FGI 2.12-3.2.2.1)	NR	2	3	Yes (ih)	NR	MERV-A-8	NR	70–75/21–24	2	10 (5) / 3	0.12 / (0.6)
Physical Therapy Exercise Area (FGI 2.12-3.2.3)	NR	2	3	Yes (ih)	NR	MERV-A-8	NR	70–75/21–24	2	20 (10) / 2	0.18 / (0.9)
Hydrotherapy (FGI 2.12-3.2.4)	Negative	2	3	Yes	NR	MERV-A-8	NR	72–80/22–27	3	20 (10) / 2	0.12 / (0.6)
Physical Therapeutic Pool (FGI 2.12-3.2.4)	Negative	2	10	Yes	NR	MERV-A-8	NR	72–80/22–27	3	----	0.48 / (2.4)
Speech Therapy Room (FGI 2.12-3.3.2)	NR	2	2	NR	NR	MERV-A-8	NR	70–75/21–24	1	5 (2.5) / 2	0.06 / (0.3)
Occupational therapy (FGI 2.12-3.3)	NR	2	3	NR	NR	MERV-A-8	NR	70–75/21–24	1	5 (2.5) / 2	0.06 / (0.3)
Prosthetics and Orthotics Room (FGI 2.12-3.3.1)	NR	2	3	NR	NR	MERV-A-8	NR	70–75/21–24	2	10 (5) / 3	0.18 / (0.9)
Dental Treatment (FGI 2.14-3.1.1)	NR	2	3	NR	NR	MERV-A-8	NR	70–75/21–24	1	10 (5) / 3	0.18 / (0.9)
Other Dental Treatment Areas (FGI 2.14-3.2)	NR	2	3	NR	NR	MERV-A-8	NR	70–75/21–24	1	5 (2.5) / 2	0.06 / (0.3)
Toilet room (FGI 2.1-3.10.2)	Negative	NR	4	Yes	No	MERV-A-8	NR	NR	3	----	----
SERVICE / SUPPORT SPACE											
Environmental services room (FGI 2.1-5.3.1)	Negative	NR	6	Yes	No	MERV-A-8	NR	NR	3	----	----

<u>Clean supply (im) (in) (FGI 2.1-3.8.11)</u>	<u>Positive</u>	<u>2</u>	<u>2</u>	<u>NR</u>	<u>NR</u>	<u>MERV-A-8</u>	<u>NR</u>	<u>NR</u>	<u>1</u>	<u>5 (2.5) / 2</u>	<u>0.12 / (0.6)</u>
<u>Soiled holding (im) (io) (ip) (FGI 2.1-3.8.12)</u>	<u>Negative</u>	<u>2</u>	<u>6</u>	<u>Yes</u>	<u>No</u>	<u>MERV-A-8</u>	<u>NR</u>	<u>NR</u>	<u>3</u>	<u>5 (2.5) / 2</u>	<u>0.12 / (0.6)</u>

Note: NR = no requirement

Normative Notes for Table 8.2:

The normative notes for Table 8.2 all begin with the letter ‘i’ (such as ia, ib, ic, etc) this is intended to differentiate from Table 8.1 yet provide some consistency in the process.

ia. Except where indicated by a “No” in this column, recirculating room HVAC units (with heating or cooling coils) are acceptable for providing that portion of the minimum total air changes per hour that is permitted by Section 8.1 (subparagraph [a][5]). Because of the cleaning difficulty and potential for buildup of contamination, recirculating room units shall not be used in areas marked “No.” Recirculating devices with high-efficiency particulate air (HEPA) filters shall be permitted in existing facilities as interim, supplemental environmental controls to meet requirements for the control of airborne infectious agents. The design of either portable or fixed systems should prevent stagnation and short circuiting of airflow. The design of such systems shall also allow for easy access for scheduled preventative maintenance and cleaning.

ib. Examination rooms (Identified as Specialty Infection Control-IC Exam rooms) programmed for use by patients with undiagnosed gastrointestinal symptoms, undiagnosed respiratory symptoms, or undiagnosed skin symptoms.

ic. Table entries are the minimum filter efficiencies required for the space. Refer to Section 6.4 of this document for further clarification of filtration requirements. Minimum filter requirements shall meet the equivalent MERV-A rating when tested in accordance with Appendix J of ANSI/ASHRAE Standard 52.2.

id. Pressure-monitoring devices are not required for any spaces indicated in Table 8.2. Simple visual methods such as ball-in-tube or flutterstrip are suggested should pressurization verification of air-flow direction be desired or required by others.

ie. Treatment rooms used for procedures with nitrous oxide shall contain provisions for exhausting anesthetic waste gases and the minimum total air changes shall be increased to 6. When this condition occurs the Rp-Ra-Air Class design option cannot be used.

if. Parenthetical notations following a space name are paragraph references to the 2018 Facility Guidelines Institute document *Guidelines for Design and Construction of Hospitals and Outpatient Facilities (Informative Note: FGI [2018])*. These FGI paragraph references are provided to the user of the standard to aid in the application of design requirements.

ig. Refer to Table 8.1 under the space name, Diagnostic imaging waiting area, for space ventilation requirements in an instance where a facility proposes an imaging waiting area and their ICRA determines that the imaging waiting area requires special consideration to reduce the risk of airborne infection transmission.

ih. In some areas with potential contamination and/or odor concerns, exhaust air shall be discharged to the outdoors and not recirculated to other areas. This exhaust may be localized by switch, timer or otherwise under operational control of the Facility Staff or Patient.

ii. The RH ranges listed are the minimum and/or maximum allowable at any point within the design temperature range required for that space.

ij. Unless otherwise noted via footnote ih, the exhaust rate shall meet or exceed minimum total air change requirement.

ik. Systems shall be capable of maintaining the rooms within the range during normal operation. Lower or higher temperature shall be permitted when patients’ comfort and/or medical conditions require those conditions.

il. This room is intended for basic blood & urine specimen testing and short term storage only. Outpatient facilities which provide the following specialized service spaces/rooms should consult table 7.1 for ventilation requirements.

1. Laboratory work areas including the specialty lab services such as bacteriology, biochemistry, cytology, glasswashing, histology, media transfer, microbiology, nuclear medicine, pathology, serology and/or sterilizing.

im. Outpatient facilities which provide the following specialized service spaces/rooms should consult table 8.1 for ventilation requirements.

1. Laundry, general and soiled linen sorting and storage

in. Refer to table 8.1 under the space name, Clean workroom or clean supply, for space ventilation requirements if a Clean workroom space is provided.

io. Refer to table 8.1 under the space name, Soiled workroom or soiled holding, for space ventilation requirements if a Soiled workroom space is provided.

ip. This space is permitted to include hazardous material storage (general medical waste). If actual regulated waste holding is anticipated refer to Table 8.1 under the space name. Regulated waste holding, of this standard for ventilation requirements.

iq. See note 6, 8 above.

8.2.3 Additional Room-Specific Requirements

~~8.2.1~~ **8.3.1 Airborne Infection Isolation (AII) Rooms.** Refer to section 7.2.1 of this standard. Ventilation for AII rooms shall meet the following requirements whenever an infectious patient occupies the room:

~~a~~ Each AII room shall comply with requirements of Tables 6.4, 6.7.2, and 8.1. AII rooms shall have a permanently installed device and/or mechanism to constantly monitor the differential air pressure between the room (when occupied by patients with a suspected airborne infectious disease) and the corridor, whether or not there is an anteroom. A local visual means shall be provided to indicate whenever negative differential pressure is not maintained.

~~b~~ All air from the AII room shall be exhausted directly to the outdoors.

Exception to 8.2.1(b): AII rooms that are retrofitted from standard patient rooms from which it is impractical to exhaust directly outdoors may be provided with recirculated air from the room's exhaust on the condition that the air first passes through a HEPA filter.

~~e~~ All exhaust air from the AII rooms, associated anterooms, and associated toilet rooms shall be discharged directly to the outdoors without mixing with exhaust air from any other non-AII room or exhaust system.

~~d~~ Exhaust air grilles or registers in the patient room shall be located directly above the patient bed, on the ceiling or on the wall near the head of the bed, unless it can be demonstrated that such a location is not practical.

~~e~~ The room envelope shall be sealed to provide a minimum differential pressure of 0.01 in. of water (2.5 Pa) across the envelope.

~~f~~ Differential pressure between AII rooms and adjacent spaces that are not AII rooms shall be a minimum of 0.01 in. of water (2.5 Pa). Spaces such as the toilet room and the anteroom (if present) that are directly associated with the AII room and open directly into the AII room are not required to be designed with a minimum pressure difference from the AII room but are still required to maintain the pressure relationships to adjacent areas specified in Table 8.1.

~~g~~ When an anteroom is provided, the pressure relationships shall be as follows: (1) the AII room shall be at a negative pressure with respect to the anteroom, and (2) the ante room shall be at a negative pressure with respect to the corridor.

8.3.2 Protective Environment (PE) Rooms. Refer to section 7.2.2 of this standard.

8.3.4 Surgery Rooms.

~~8.3.1~~ **8.4.1 Operating Rooms, Operating/Surgical Cystoscopic Rooms, and Caesarean Delivery Rooms.**

Refer to section 7.4.1 of this standard. These rooms shall be maintained at a positive pressure with respect to all adjoining spaces at all times. A pressure differential shall be maintained at a value of at least +0.01 in. wc (2.5 Pa). Each room shall have individual temperature control. These rooms shall be provided with a primary supply diffuser array that is designed as follows:

~~a~~ The airflow shall be unidirectional, downwards, and the average velocity of the diffusers shall be 25 to 35 cfm/ft² (127 to 178 L/s/m²). The diffusers shall be concentrated to provide an airflow pattern over the patient and surgical team. *Informative Note:* For more information, see Memarzadeh and Manning (2002) and Memarzadeh and Jiang (2004) in Appendix B.

~~b~~ The coverage area of the primary supply diffuser array shall extend a minimum of 12 in. (305 mm) beyond the footprint of the surgical table on each side. Within the portion of the primary supply diffuser array that consists of an area encompassing 12 in. (305 mm) on each side of the footprint of the surgical table, no more than 30% of this portion of the primary supply diffuser array area shall be used for nondiffuser uses such as lights, gas columns, equipment booms, access panels, sprinklers, etc.

Additional supply diffusers shall be permitted within the room, outside of the primary supply diffuser array, to provide additional ventilation to the operating room to achieve the environmental requirements of Table 8.1 that relate to temperature, humidity, or a portion of the required air change rates.

~~The room shall be provided with at least two low side wall return or exhaust grilles spaced at opposite corners or as far apart as possible, with the bottom of these grilles installed approximately 8 in. (203 mm) above the floor.~~

~~**Exception to 8.3.1:** In addition to the required low return (or exhaust) air grilles, such grilles may be placed high on the walls.~~

~~**8.3.2 8.4.2 Sterilization Rooms.** Steam that escapes from a steam sterilizer shall be exhausted using an exhaust hood or other suitable means. Ethylene oxide that escapes from a gas sterilizer shall be exhausted using an exhaust hood or other suitable means.~~

~~**8.3.3 8.4.3 Imaging Procedure Rooms.** If invasive procedures occur in this type of room, ventilation shall be provided in accordance with the ventilation requirements for procedure rooms. If anesthetic gases are administered, ventilation shall be provided in accordance with the ventilation requirements for operating rooms.~~

8.4 8.5 Support Spaces.

~~**8.4.1 8.5.1 Nonrefrigerated Body-Holding Rooms.** This space type is not listed in Tables 8.1 or 8.2, however, the following specific functional elements are relative to any Facility with a Body-Holding Room. A nonrefrigerated body-holding room is applicable only to facilities that do not perform autopsies on-site and use the space for short periods while waiting for the body to be transferred. Ventilation for nonrefrigerated body-holding rooms shall meet the following requirements:~~

~~a. All exhaust air from nonrefrigerated body holding rooms shall be discharged directly to the outdoors without mixing with air from any other room or exhaust system.~~

~~**8.4.2 8.5.2 Bronchoscopy.**~~

~~a. Differential pressure between bronchoscopy procedure and sputum induction rooms and any adjacent spaces that have other functions shall be a minimum of -0.01 in. of water (-2.5 Pa).~~

~~b. Local exhaust shall be provided for sputum collection procedures.~~

~~**8.5 8.6 Psychiatric Patient Areas.** All exposed equipment located with these spaces shall have enclosures with rounded corners and tamper-resistant fasteners. With the exception of HVAC room recirculating units, equipment shall be arranged such that maintenance personnel are not required to enter patient-care spaces for service.~~

~~**8.7 Nuclear Medicine.** Refer to Table 8.1 of this standard for both nuclear medicine treatment spaces and nuclear medicine hot lab spaces when radiopharmaceutical preparation is performed on-site (not pre-mixed) and radioactive materials (radionuclides) are mixed/distributed from their protective containers within this room. If dose administration and on-site mixing and preparation is using only low level pre-mixed radioactive materials then a 'hot lab' is not indicated and these nuclear medicine spaces will follow the general examination room space in Table 8.2 of this standard for ventilation requirements.~~