





*Protecting, Maintaining and Improving the Health of All Minnesotans*

Electronically delivered  
October 10, 2021

Administrator  
Minneota Manor Health Care Center  
700 North Monroe Street  
Minneota, MN 56264

RE: CCN: 245496  
Cycle Start Date: September 16, 2021

Dear Administrator:

On September 16, 2021, a survey was completed at your facility by the Minnesota Departments of Health and Public Safety, to determine if your facility was in compliance with Federal participation requirements for skilled nursing facilities and/or nursing facilities participating in the Medicare and/or Medicaid programs.

This survey found the most serious deficiencies in your facility to be widespread deficiencies that constituted no actual harm with potential for more than minimal harm that was not immediate jeopardy (Level F), as evidenced by the electronically attached CMS-2567 whereby corrections are required.

### **ELECTRONIC PLAN OF CORRECTION (ePoC)**

Within **ten (10) calendar days** after your receipt of this notice, you must submit an acceptable ePOC for the deficiencies cited. An acceptable ePOC will serve as your allegation of compliance. Upon receipt of an acceptable ePOC, we will authorize a revisit to your facility to determine if substantial compliance has been achieved.

To be acceptable, a provider's ePOC must include the following:

- How corrective action will be accomplished for those residents found to have been affected by the deficient practice.
- How the facility will identify other residents having the potential to be affected by the same deficient practice.
- What measures will be put into place, or systemic changes made, to ensure that the deficient practice will not recur.
- How the facility will monitor its corrective actions to ensure that the deficient practice is being corrected and will not recur.
- The date that each deficiency will be corrected.
- An electronic acknowledgement signature and date by an official facility representative.

The state agency may, in lieu of an onsite revisit, determine correction and compliance by accepting the facility's ePoC if the ePoC is reasonable, addresses the problem and provides evidence that the corrective action has occurred.

If an acceptable ePoC is not received within 10 calendar days from the receipt of this letter, we will recommend to the CMS Region V Office that one or more of the following remedies be imposed:

- Denial of payment for new Medicare and Medicaid admissions (42 CFR 488.417);
- Civil money penalty (42 CFR 488.430 through 488.444).
- Termination of your facility's Medicare and/or Medicaid agreement (488.456(b)).

## DEPARTMENT CONTACT

Questions regarding this letter and all documents submitted as a response to the resident care deficiencies (those preceded by an "F" tag) and emergency preparedness deficiencies (those preceded by an "E" tag), i.e., the plan of correction should be directed to:

**Nicole Osterloh, RN, Unit Supervisor**  
**Marshall District Office**  
**Licensing and Certification Program**  
**Health Regulation Division**  
**Minnesota Department of Health**  
**1400 East Lyon Street, Suite 102**  
**Marshall, Minnesota 56258-2504**  
**Email: nicole.osterloh@state.mn.us**  
**Office: 507-476-4230**  
**Mobile: (507) 251-6264 Mobile: (605) 881-6192**

## PRESUMPTION OF COMPLIANCE - CREDIBLE ALLEGATION OF COMPLIANCE

The facility's ePoC will serve as your allegation of compliance upon the Department's acceptance. In order for your allegation of compliance to be acceptable to the Department, the ePoC must meet the criteria listed in the plan of correction section above. You will be notified by the Minnesota Department of Health, Licensing and Certification Program staff and/or the Department of Public Safety, State Fire Marshal Division staff, if your ePoC for the respective deficiencies (if any) is acceptable.

## VERIFICATION OF SUBSTANTIAL COMPLIANCE

Upon receipt of an acceptable ePoC, a Post Certification Revisit (PCR), of your facility will be conducted to validate that substantial compliance with the regulations has been attained in accordance with your verification.

If substantial compliance has been achieved, certification of your facility in the Medicare and/or

Medicaid program(s) will be continued and remedies will not be imposed. Compliance is certified as of the latest correction date on the approved ePoC, unless it is determined that either correction actually occurred between the latest correction date on the ePoC and the date of the first revisit, or correction occurred sooner than the latest correction date on the ePoC.

#### **FAILURE TO ACHIEVE SUBSTANTIAL COMPLIANCE BY THE THIRD OR SIXTH MONTH AFTER THE LAST DAY OF THE SURVEY**

If substantial compliance with the regulations is not verified by December 16, 2021 (three months after the identification of noncompliance), the CMS Region V Office must deny payment for new admissions as mandated by the Social Security Act (the Act) at Sections 1819(h)(2)(D) and 1919(h)(2)(C) and Federal regulations at 42 CFR Section 488.417(b).

In addition, if substantial compliance with the regulations is not verified by March 16, 2022 (six months after the identification of noncompliance) your provider agreement will be terminated. This action is mandated by the Social Security Act at Sections 1819(h)(2)(C) and 1919(h)(3)(D) and Federal regulations at 42 CFR Sections 488.412 and 488.456.

**Please note that this notice does not constitute formal notice of imposition of alternative remedies or termination of your provider agreement. Should the Centers for Medicare & Medicaid Services determine that termination or any other remedy is warranted, it will provide you with a separate formal notification of that determination.**

#### **INFORMAL DISPUTE RESOLUTION (IDR) / INDEPENDENT INFORMAL DISPUTE RESOLUTION (IIDR)**

In accordance with 42 CFR 488.331, you have one opportunity to question cited deficiencies through an informal dispute resolution process. You are required to send your written request, along with the specific deficiencies being disputed, and an explanation of why you are disputing those deficiencies, to:

Nursing Home Informal Dispute Process  
Minnesota Department of Health  
Health Regulation Division  
P.O. Box 64900  
St. Paul, Minnesota 55164-0900

This request must be sent within the same ten days you have for submitting an ePoC for the cited deficiencies. All requests for an IDR or IIDR of federal deficiencies must be submitted via the web at: [https://mdhprovidercontent.web.health.state.mn.us/ltc\\_idr.cfm](https://mdhprovidercontent.web.health.state.mn.us/ltc_idr.cfm)

You must notify MDH at this website of your request for an IDR or IIDR within the 10 calendar day period allotted for submitting an acceptable electronic plan of correction. A copy of the Department's informal dispute resolution policies are posted on the MDH Information Bulletin website at: [https://www.health.state.mn.us/facilities/regulation/infobulletins/ib04\\_8.html](https://www.health.state.mn.us/facilities/regulation/infobulletins/ib04_8.html)

Please note that the failure to complete the informal dispute resolution process will not delay the dates

Minneota Manor Health Care Center

October 10, 2021

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specified for compliance or the imposition of remedies.

Questions regarding all documents submitted as a response to the Life Safety Code deficiencies (those preceded by a "K" tag), i.e., the plan of correction, request for waivers, should be directed to:

**William Abderhalden, Fire Safety Supervisor**  
**Deputy State Fire Marshal**  
**Health Care/Corrections Supervisor – Interim**  
**Minnesota Department of Public Safety**  
**445 Minnesota Street, Suite 145**  
**St. Paul, MN 55101-5145**  
**Cell: (507) 361-6204**  
**Email: [william.abderhalden@state.mn.us](mailto:william.abderhalden@state.mn.us)**  
**Fax: (651) 215-0525**

Feel free to contact me if you have questions.

Sincerely,



Kamala Fiske-Downing  
Minnesota Department of Health  
Licensing and Certification Program  
Program Assurance Unit  
Health Regulation Division  
Telephone: (651) 201-4112 Fax: (651) 215-9697  
Email: [Kamala.Fiske-Downing@state.mn.us](mailto:Kamala.Fiske-Downing@state.mn.us)



*Protecting, Maintaining and Improving the Health of All Minnesotans*

Electronically delivered  
November 24, 2021

Administrator  
Minneota Manor Health Care Center  
700 North Monroe Street  
Minneota, MN 56264

RE: CCN: 245496  
Cycle Start Date: September 16, 2021

Dear Administrator:

On October 10, 2021, we informed you that we may impose enforcement remedies.

Compliance with the Life Safety Code (LSC) deficiencies cited on September 16, 2021 has not yet been verified.

Sections 1819(h)(2)(D) and (E) and 1919(h)(2)(C) and (D) of the Act and 42 CFR 488.417(b) require that, regardless of any other remedies that may be imposed, denial of payment for new admissions must be imposed when the facility is not in substantial compliance 3 months after the last day of the survey identifying noncompliance. Thus, the CMS Region V Office concurs, is imposing the following remedy and has authorized this Department to notify you of the imposition:

- Mandatory Denial of payment for new Medicare and Medicaid admissions effective September 16, 2021. (42 CFR 488.417 (b))

The CMS Region V Office will notify your Medicare Administrative Contractor (MAC) that the denial of payment for new admissions is effective September 16, 2021. They will also notify the State Medicaid Agency that they must also deny payment for new Medicaid admissions effective September 16, 2021. You should notify all Medicare/Medicaid residents admitted on or after this date of the restriction.

Further, Federal law, as specified in the Act at Sections 1819(f)(2)(B), prohibits approval of nurse assistant training programs offered by, or in, a facility which, within the previous two years, has been subject to a denial of payment. Therefore, Minneota Manor Health Care Center is prohibited from offering or conducting a Nurse Assistant Training/Competency Evaluation Programs or Competency Evaluation Programs for two years effective September 16, 2021. This prohibition is not subject to appeal. Further, this prohibition may be rescinded at a later date if your facility achieves substantial compliance prior to the effective date of denial of payment for new admissions. If this prohibition is not rescinded, under Public Law 105-15 (H.R. 968), you may request a waiver of this prohibition if certain criteria are met. Please contact the Nursing Assistant Registry at (800) 397-6124 for specific information regarding a waiver for these programs from this Department.

## **APPEAL RIGHTS**

If you disagree with this action imposed on your facility, you or your legal representative may request a hearing before an administrative law judge of the Department of Health and Human Services, Departmental Appeals Board (DAB). Procedures governing this process are set out in 42 C.F.R. 498.40, et seq. You must file your hearing request electronically by using the Departmental Appeals Board's Electronic Filing System (DAB E-File) at <https://dab.efile.hhs.gov> no later than sixty (60) days after receiving this letter. Specific instructions on how to file electronically are attached to this notice. A copy of the hearing request shall be submitted electronically to:

**Tamika.Brown@cms.hhs.gov**

Requests for a hearing submitted by U.S. mail or commercial carrier are no longer accepted as of October 1, 2014, unless you do not have access to a computer or internet service. In those circumstances you may call the Civil Remedies Division to request a waiver from e-filing and provide an explanation as to why you cannot file electronically or you may mail a written request for a waiver along with your written request for a hearing. A written request for a hearing must be filed no later than sixty (60) days after receiving this letter, by mailing to the following address:

**Department of Health & Human Services  
Departmental Appeals Board, MS 6132  
Director, Civil Remedies Division  
330 Independence Avenue, S.W.  
Cohen Building – Room G-644  
Washington, D.C. 20201  
(202) 565-9462**

A request for a hearing should identify the specific issues, findings of fact and conclusions of law with which you disagree. It should also specify the basis for contending that the findings and conclusions are incorrect. At an appeal hearing, you may be represented by counsel at your own expense. If you have any questions regarding this matter, please contact Tamika Brown, Principal Program Representative by phone at (312) 353-1502 or by e-mail at [Tamika.Brown@cms.hhs.gov](mailto:Tamika.Brown@cms.hhs.gov).

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In accordance with 42 CFR 488.331, you have one opportunity to question cited deficiencies through an informal dispute resolution process. You are required to send your written request, along with the specific deficiencies being disputed, and an explanation of why you are disputing those deficiencies, to:

**Nursing Home Informal Dispute Process  
Minnesota Department of Health  
Health Regulation Division  
P.O. Box 64900  
St. Paul, Minnesota 55164-0900**

This request must be sent within the same ten days you have for submitting an ePoC for the cited deficiencies. All requests for an IDR or IIDR of federal deficiencies must be submitted via the web at: [https://mdhprovidercontent.web.health.state.mn.us/ltc\\_idr.cfm](https://mdhprovidercontent.web.health.state.mn.us/ltc_idr.cfm)

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Please note that the failure to complete the informal dispute resolution process will not delay the dates specified for compliance or the imposition of remedies.

Questions regarding all documents submitted as a response to the Life Safety Code deficiencies (those preceded by a "K" tag) i.e., the plan of correction, request for waivers, should be directed to:

William Abderhalden, Fire Safety Supervisor  
Deputy State Fire Marshal  
Health Care/Corrections Supervisor – Interim  
Minnesota Department of Public Safety  
445 Minnesota Street, Suite 145  
St. Paul, MN 55101-5145  
Cell: (507) 361-6204  
Email: [william.abderhalden@state.mn.us](mailto:william.abderhalden@state.mn.us)  
Fax: (651) 215-0525

Feel free to contact me if you have questions.

Sincerely,



Kamala Fiske-Downing  
Minnesota Department of Health  
Licensing and Certification Program  
Program Assurance Unit  
Health Regulation Division  
Telephone: (651) 201-4112 Fax: (651) 215-9697  
Email: [Kamala.Fiske-Downing@state.mn.us](mailto:Kamala.Fiske-Downing@state.mn.us)



DEPARTMENT OF HEALTH AND HUMAN SERVICES  
CENTERS FOR MEDICARE & MEDICAID SERVICES

F5496031

PRINTED: 11/04/2021  
FORM APPROVED  
OMB NO. 0938-0391

STATEMENT OF DEFICIENCIES AND PLAN OF CORRECTION		(X1) PROVIDER/SUPPLIER/CLIA IDENTIFICATION NUMBER:  <b>245496</b>		(X2) MULTIPLE CONSTRUCTION A. BUILDING <b>01 - MAIN BUILDING 01</b>  B. WING _____		(X3) DATE SURVEY COMPLETED  <b>09/16/2021</b>	
NAME OF PROVIDER OR SUPPLIER  <b>MINNEOTA MANOR HEALTH CARE CENTER</b>				STREET ADDRESS, CITY, STATE, ZIP CODE <b>700 NORTH MONROE STREET MINNEOTA, MN 56264</b>			
(X4) ID PREFIX TAG	SUMMARY STATEMENT OF DEFICIENCIES (EACH DEFICIENCY MUST BE PRECEDED BY FULL REGULATORY OR LSC IDENTIFYING INFORMATION)			ID PREFIX TAG	PROVIDER'S PLAN OF CORRECTION (EACH CORRECTIVE ACTION SHOULD BE CROSS-REFERENCED TO THE APPROPRIATE DEFICIENCY)		(X5) COMPLETION DATE
K 000	<p>INITIAL COMMENTS</p> <p>FIRE SAFETY</p> <p>An annual Life Safety Code survey was conducted by the Minnesota Department of Public Safety, State Fire Marshal Division on 09/15/2021. At the time of this survey, MINNEOTA MANOR HEALTH CARE CENTER was found not in compliance with the requirements for participation in Medicare/Medicaid at 42 CFR, Subpart 483.70(a), Life Safety from Fire, and the 2012 edition of National Fire Protection Association (NFPA) 101, Life Safety Code (LSC), Chapter 19 Existing Health Care and the 2012 edition of NFPA 99, Health Care Facilities Code.</p> <p>THE FACILITY'S POC WILL SERVE AS YOUR ALLEGATION OF COMPLIANCE UPON THE DEPARTMENT'S ACCEPTANCE. YOUR SIGNATURE AT THE BOTTOM OF THE FIRST PAGE OF THE CMS-2567 FORM WILL BE USED AS VERIFICATION OF COMPLIANCE.</p> <p>UPON RECEIPT OF AN ACCEPTABLE POC, AN ONSITE REVISIT OF YOUR FACILITY MAY BE CONDUCTED TO VALIDATE THAT SUBSTANTIAL COMPLIANCE WITH THE REGULATIONS HAS BEEN ATTAINED IN ACCORDANCE WITH YOUR VERIFICATION.</p> <p>PLEASE RETURN THE PLAN OF CORRECTION FOR THE FIRE SAFETY DEFICIENCIES (K-TAGS) TO:</p> <p>IF PARTICIPATING IN THE E-POC PROCESS, A PAPER COPY OF THE PLAN OF CORRECTION IS NOT REQUIRED.</p>			K 000			

LABORATORY DIRECTOR'S OR PROVIDER/SUPPLIER REPRESENTATIVE'S SIGNATURE

TITLE

(X6) DATE

Electronically Signed

10/20/2021

Any deficiency statement ending with an asterisk (\*) denotes a deficiency which the institution may be excused from correcting providing it is determined that other safeguards provide sufficient protection to the patients. (See instructions.) Except for nursing homes, the findings stated above are disclosable 90 days following the date of survey whether or not a plan of correction is provided. For nursing homes, the above findings and plans of correction are disclosable 14 days following the date these documents are made available to the facility. If deficiencies are cited, an approved plan of correction is requisite to continued program participation.

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NAME OF PROVIDER OR SUPPLIER  <b>MINNEOTA MANOR HEALTH CARE CENTER</b>			STREET ADDRESS, CITY, STATE, ZIP CODE <b>700 NORTH MONROE STREET MINNEOTA, MN 56264</b>		
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K 000	<p>Continued From page 1</p> <p>Healthcare Fire Inspections State Fire Marshal Division 445 Minnesota St., Suite 145 St. Paul, MN 55101-5145, OR</p> <p>By email to: FM.HC.Inspections@state.mn.us</p> <p>THE PLAN OF CORRECTION FOR EACH DEFICIENCY MUST INCLUDE ALL OF THE FOLLOWING INFORMATION:</p> <ol style="list-style-type: none"> <li>1. A detailed description of the corrective action taken or planned to correct the deficiency.</li> <li>2. Address the measures that will be put in place to ensure the deficiency does not reoccur.</li> <li>3. Indicate how the facility plans to monitor future performance to ensure solutions are sustained.</li> <li>4. Identify who is responsible for the corrective actions and monitoring of compliance.</li> <li>5. The actual or proposed date for completion of the remedy.</li> </ol> <p>MINNEOTA MANOR HEALTH CARE CENTER was constructed at two different times. A one-story building with no basement was constructed in 1972 and determined to be Type II ( 111 ). In 1995, a one-story building with no basement was constructed and determined to be Type II (111)</p>	K 000			

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K 000	Continued From page 2 Because the original building and additions are compatible construction types allowed for existing buildings of this height, the facility was surveyed as one building as allowed in the 2012 edition of National Fire Protection Association (NFPA) Standard 101, Life Safety Code (LSC), Chapter 19 Existing Health Care Occupancies.  The facility is fully protected throughout by an automatic sprinkler system and has a fire alarm system with smoke detection in the corridors, spaces open to the corridors that is monitored for automatic fire department notification.  The facility has a capacity of 55 beds and had a census of 23 at the time of the survey.	K 000			
K 271 SS=E	The requirement at 42 CFR, Subpart 483.70(a) is NOT MET as evidence by: Discharge from Exits CFR(s): NFPA 101  Discharge from Exits Exit discharge is arranged in accordance with 7.7, provides a level walking surface meeting the provisions of 7.1.7 with respect to changes in elevation and shall be maintained free of obstructions. Additionally, the exit discharge shall be a hard packed all-weather travel surface. 18.2.7, 19.2.7 This REQUIREMENT is not met as evidenced by: Based on observation and staff interview, the facility failed to maintain the exit discharge in accordance with the NFPA 101 (2012 edition), Life Safety Code, sections 19.2.7, 7.1.6.2, 7.1.7, 7.7. This deficient condition could have a patterned impact on the residents within the	K 271	K271 I. Corrective Action Taken  1. South Exit Door -Cement will be ground down to grade 12-3-21 2. North Exit Door <input type="checkbox"/> Replaced with new		12/3/21

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NAME OF PROVIDER OR SUPPLIER  <b>MINNEOTA MANOR HEALTH CARE CENTER</b>			STREET ADDRESS, CITY, STATE, ZIP CODE <b>700 NORTH MONROE STREET MINNEOTA, MN 56264</b>		
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K 271	Continued From page 3 facility.  Findings include:  1. On 09/16/2021 between 11:30 AM to 04:30 PM, it was revealed during the walk-through of the facility that the South Exit Door ( Wing 200 ) egress to grade had a vertical displacement greater than one-half inch, and the concrete was degraded and breaking up, presenting a fall and trip hazard.  2. On 09/16/2021 between 11:30 AM to 04:30 PM, it was revealed during the walk-through of the facility that the North Exit Door ( Wing 100 ) egress to grade had a horizontal separation of 2-inches between concrete slabs breaking up, presenting a fall and trip hazard.  This deficient condition was confirmed by the Facility Maintenance Director at the time of discovery.	K 271	cement 10-1-21  II. Measures put in place to ensure it does not reoccur Facility will no longer be operating due to facility closure 12-3-21 III. Monitoring Facility will no longer be operating due to facility closure 12-3-21  IV. Who is responsible Maintenance Director V. Actual date for completion of the remedy 12-3-21		
K 353 SS=F	Sprinkler System - Maintenance and Testing CFR(s): NFPA 101  Sprinkler System - Maintenance and Testing Automatic sprinkler and standpipe systems are inspected, tested, and maintained in accordance with NFPA 25, Standard for the Inspection, Testing, and Maintaining of Water-based Fire Protection Systems. Records of system design, maintenance, inspection and testing are maintained in a secure location and readily available. a) Date sprinkler system last checked  b) Who provided system test	K 353		12/3/21	

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K 353	<p>Continued From page 4</p> <p>c) Water system supply source</p> <p>Provide in REMARKS information on coverage for any non-required or partial automatic sprinkler system. 9.7.5, 9.7.7, 9.7.8, and NFPA 25 This REQUIREMENT is not met as evidenced by: Based on observation and staff interview, the facility failed to inspect and maintain the sprinkler system in accordance with NFPA 101 (2012 edition), Life Safety Code, sections 9.7.5, 9.7.6, and NFPA 25 (2011 edition) Standard for the Inspection, Testing, and Maintenance of Water-Based Fire Protection Systems, sections 5.2, 5.2.1.1.1, 5.2.1.1.2, 5.2.1.1.4, 5.2.1.2. NFPA13 (2010 edition), Standard for the Installation of Sprinkler Systems, sections 8.5.6, 8.5.6.1. This deficient condition could have a widespread impact on the residents within the facility.</p> <p>Findings include:</p> <p>1. On 09/16/2021 between 11:30 AM to 04:30 PM, it was revealed during the walk-through of the facility that oxidized sprinkler heads were observed in the Kitchen - stove area and dish-washing area.</p> <p>2. On 09/16/2021 between 11:30 AM to 04:30 PM, it was revealed during the walk-through of the facility that items were stacked too high, less than 18 inches, in the following areas: S6, RM 205, N3, N4, N5, Linen closet adjacent to RM 209, storage closet adjacent to RM 209.</p> <p>3. On 09/16/2021 between 11:30 AM to 04:30 PM, it was revealed during the walk-through of</p>	K 353	<p>K353</p> <p>I. Corrective Action Taken</p> <p>1. Oxidized sprinkler heads in kitchen-stove area replaced 10-20-21</p> <p>2. S6, RM205, N#, N4, N5, linen closet adjacent to RM209, storage closet adjacent to RM209 □ items stacked too high: Removed items and removed top shelf. Closets were marked with red tape to indicate the maximum storage height. 9-28-21</p> <p>3. 5 year inspection of sprinkler system. Summit scheduled to complete 5 year inspection of the sprinkler system on 10-20-21. Completion date 12-3-21.</p> <p>II. Measures put in place to ensure deficient practice does not reoccur: Facility will no longer be operating due to facility closure 12-3-21</p> <p>III. Monitoring Facility will no longer be operating due to facility closure 12-3-21</p> <p>IV. Who is responsible Maintenance Director</p> <p>V. Actual date for completions of remedy : 12-3-21</p>		

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K 353	Continued From page 5 the facility and documentation review that the most recent 5-year inspection of the sprinkler system was complete on 07/16/2014.	K 353			
K 374 SS=F	<p>This deficient condition was confirmed by the Facility Maintenance Director at the time of discovery.</p> <p>Subdivision of Building Spaces - Smoke Barrie CFR(s): NFPA 101</p> <p>Subdivision of Building Spaces - Smoke Barrier Doors 2012 EXISTING Doors in smoke barriers are 1-3/4-inch thick solid bonded wood-core doors or of construction that resists fire for 20 minutes. Nonrated protective plates of unlimited height are permitted. Doors are permitted to have fixed fire window assemblies per 8.5. Doors are self-closing or automatic-closing, do not require latching, and are not required to swing in the direction of egress travel. Door opening provides a minimum clear width of 32 inches for swinging or horizontal doors. 19.3.7.6, 19.3.7.8, 19.3.7.9 This REQUIREMENT is not met as evidenced by: Based on observation and staff interview, the facility failed to inspect and maintain smoke barrier doors per NFPA 101 (2012 edition), Life Safety Code, sections 19.3.7 and 8.5.4. These deficient conditions could have a widespread impact on the residents within the facility.</p> <p>Findings include:</p> <p>On 09/16/2021 between 11:30 AM to 04:30 PM, it was revealed during the walk-through of the</p>	K 374	<p>K374</p> <p>I. Corrective Action Taken</p> <p>1. Smoke barrier doors adjacent to RM W 306 and RM W 309 Doors were adjusted so they sealed correctly 10-5-21</p> <p>II. Measures put in place to ensure deficient practice does not reoccur: Facility will no longer be operating due to</p>	12/3/21	

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K 374	Continued From page 6 facility that upon testing of the smoke barrier doors adjacent to RM W-306 and those adjacent to RM W-309, the doors did not close and seal properly.  These deficient conditions were confirmed by the Maintenance Director at the time of discovery.	K 374	facility closure 12-3-21  III. Monitoring Facility will no longer be operating due to facility closure 12-3-21  IV. Who is responsible Maintenance Director V. Actual date for completions of remedy 12-3-21		
K 511 SS=F	Utilities - Gas and Electric CFR(s): NFPA 101  Utilities - Gas and Electric Equipment using gas or related gas piping complies with NFPA 54, National Fuel Gas Code, electrical wiring and equipment complies with NFPA 70, National Electric Code. Existing installations can continue in service provided no hazard to life. 18.5.1.1, 19.5.1.1, 9.1.1, 9.1.2  This REQUIREMENT is not met as evidenced by: Based on observation and staff interview, the facility failed to maintain security and accessibility to electrical panels in a resident accessible corridor in accordance with NFPA 101 (2012 edition), Life Safety Code, sections 19.5.1.1 and 9.1.2, NFPA 70 (2011 edition), National Electrical Code, section 110.26, and NFPA 99, (2012 edition), Health Care Facilities Code, section 6.3.2.2.1.3. These deficient conditions could have a widespread impact on the residents within the facility.	K 511	K511 I. Corrective Action 1. South wing electric panel unsecured: Paddle locks installed 10-4-21 2. Dining room area adjacent to RM W316 □ Paddle locks installed 10-4-21  II. Measures put in place to ensure deficient practice does not reoccur: Facility will no longer be operating due to facility closure 12-3-21		12/3/21

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K 511	Continued From page 7  Findings include:  On 09/16/2021 between 11:30 AM to 04:30 PM, it was revealed during the walk-through of the facility that the following electrical panels in resident accessible corridors were unsecured: 1. South Wing - panel by Nurses Station 2. Dining Room area - adjacent to RM W-316  These deficient conditions were confirmed by the Maintenance Director at the time of discovery.	K 511	III. Monitoring Facility will no longer be operating due to facility closure 12-3-21 IV. Who is responsible Maintenance Director V. Actual date for completions of remedy 12-3-21		
K 521 SS=F	HVAC CFR(s): NFPA 101  HVAC Heating, ventilation, and air conditioning shall comply with 9.2 and shall be installed in accordance with the manufacturer's specifications. 18.5.2.1, 19.5.2.1, 9.2  This REQUIREMENT is not met as evidenced by: Based on observations, documentation review, and staff interview, it was revealed that the facility is using the corridors as an air plenum which is not in accordance with NFPA 101 (2012 edition), Life Safety Code, section 19.5.2, and NFPA 90A (2012 edition), Standard for the Installation of Air-Conditioning and Ventilating Systems, section 4.3.12.1.1. This deficient condition could have a widespread impact on the residents within the facility.	K 521	K521 Application for Federal Waiver attached	12/3/21	



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K 521	Continued From page 8 Findings include:  On 09/16/2021 between 11:30 AM to 04:30 PM, observations, documentation review, and staff interview revealed the ventilation system in the 1972 building utilized the egress corridors as a return air plenum for the building HVAC system. Specifically, resident rooms were equipped with supply air diffusers only, and the bathroom exhaust fans were switched, i.e., did not run continuously. Further, the concealed spaces above the drop-ceiling assembly in the egress corridors were used to provide the return air for the building HVAC system.	K 521			
K 712 SS=F	This deficient condition was verified by the Maintenance Supervisor.  Fire Drills CFR(s): NFPA 101  Fire Drills Fire drills include the transmission of a fire alarm signal and simulation of emergency fire conditions. Fire drills are held at expected and unexpected times under varying conditions, at least quarterly on each shift. The staff is familiar with procedures and is aware that drills are part of established routine. Where drills are conducted between 9:00 PM and 6:00 AM, a coded announcement may be used instead of audible alarms. 19.7.1.4 through 19.7.1.7 This REQUIREMENT is not met as evidenced by: Based on document review and staff interview, the facility failed to conduct fire drills in accordance with the NFPA 101 (2012 edition), Life Safety Code, sections 19.7.1.6, 4.7.2, and	K 712			12/3/21
			K712 I. Corrective action taken Reviewed drills 10-1-21 to ensure the correct rotation was being followed.		

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K 712	Continued From page 9 4.7.6. This deficient condition could have a widespread impact on the residents within the facility.  Findings include:  On 09/16/2021 between 11:30 AM to 04:30 PM, it was revealed during documentation review that no documentation was presented for review to confirm that fire drills were conducted for 2nd shift staff in the 1st and 2nd quarters.  This deficient condition was confirmed by the Facility Maintenance Director at the time of discovery.	K 712	II. To ensure deficient practice does not reoccur, Maintenance Director will Monitor November's fire drill to assure it is on the correct shift. III. Monitoring Maintenance Director will monitor November's fire drill to assure it is on the correct shift. IV. Who is responsible: Maintenance Director V. Date of completion: 12-3-21		
K 920 SS=F	Electrical Equipment - Power Cords and Extens CFR(s): NFPA 101  Electrical Equipment - Power Cords and Extension Cords Power strips in a patient care vicinity are only used for components of movable patient-care-related electrical equipment (PCREE) assemblies that have been assembled by qualified personnel and meet the conditions of 10.2.3.6. Power strips in the patient care vicinity may not be used for non-PCREE (e.g., personal electronics), except in long-term care resident rooms that do not use PCREE. Power strips for PCREE meet UL 1363A or UL 60601-1. Power strips for non-PCREE in the patient care rooms (outside of vicinity) meet UL 1363. In non-patient care rooms, power strips meet other UL standards. All power strips are used with general precautions. Extension cords are not used as a substitute for fixed wiring of a structure. Extension cords used temporarily are removed immediately upon completion of the purpose for	K 920		12/3/21	

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K 920	<p>Continued From page 10</p> <p>which it was installed and meets the conditions of 10.2.4.</p> <p>10.2.3.6 (NFPA 99), 10.2.4 (NFPA 99), 400-8 (NFPA 70), 590.3(D) (NFPA 70), TIA 12-5</p> <p>This REQUIREMENT is not met as evidenced by:</p> <p>Based on observation and staff interview, the facility failed to implement the usage of power strips in accordance with NFPA 99 (2012 edition), Health Care Facilities Code, section 10.2.3.6, 10.2.4 and NFPA 70, (2011 edition), National Electrical Code, sections 400-8, 590.3(D). These deficient conditions could have a widespread impact on the residents within the facility.</p> <p>Findings include:</p> <p>1. On 09/16/2021 between 11:30 AM to 04:30 PM, it was revealed during the facility walk-through of the facility that appliances were connected to power-strips in the following rooms: C-5 and W-2.</p> <p>2. On 09/16/2021 between 11:30 AM to 04:30 PM, it was revealed during the facility walk-through of the facility that an extension cord was in use in the North Day Room.</p> <p>3. On 09/16/2021 between 11:30 AM to 04:30 PM, it was revealed during the facility walk-through of the facility that a 2-to-6 multi-tap electrical outlet adapter was in use in the Social Services Office.</p> <p>4. On 09/16/2021 between 11:30 AM to 04:30 PM, it was revealed during the facility walk-through of the facility that two triple-tap electrical outlet adapters were in use in a resident room ( RM 104 ).</p>	K 920	<p>K920</p> <p>I. Corrective Action taken</p> <p>1. Appliances connected to power-strips in rooms C5 &amp; W2: Removed power strips 9-21-21</p> <p>2. Extension cord was in use in N dayroom: Removed extension cord 9-21-21</p> <p>3. 2-6 multi-tap electrical outlet adapter was in use in SS office: removed 9-21-21</p> <p>4. Two triple electrical outlet adapters were in use in a RM 104: removed adapters 9-21-21</p> <p>II. Ensure deficient practice does not reoccur:</p> <p>Maintenance Director will conduct walk through <input type="checkbox"/> admissions are not allowed during closure, so no new hook ups are taking place at this time.</p> <p>III. Maintenance Director will walk through and observe until closure 12-3-21</p> <p>IV. Who is responsible <input type="checkbox"/> Maintenance Director</p> <p>V. Date of Completion: 12-3-21</p>		

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K 920	Continued From page 11			K 920			
K 923 SS=F	<p>These deficient conditions were confirmed by the Maintenance Director at the time of discovery.</p> <p>Gas Equipment - Cylinder and Container Storage CFR(s): NFPA 101</p> <p>Gas Equipment - Cylinder and Container Storage Greater than or equal to 3,000 cubic feet Storage locations are designed, constructed, and ventilated in accordance with 5.1.3.3.2 and 5.1.3.3.3. &gt;300 but &lt;3,000 cubic feet Storage locations are outdoors in an enclosure or within an enclosed interior space of non- or limited- combustible construction, with door (or gates outdoors) that can be secured. Oxidizing gases are not stored with flammables, and are separated from combustibles by 20 feet (5 feet if sprinklered) or enclosed in a cabinet of noncombustible construction having a minimum 1/2 hr. fire protection rating. Less than or equal to 300 cubic feet In a single smoke compartment, individual cylinders available for immediate use in patient care areas with an aggregate volume of less than or equal to 300 cubic feet are not required to be stored in an enclosure. Cylinders must be handled with precautions as specified in 11.6.2. A precautionary sign readable from 5 feet is on each door or gate of a cylinder storage room, where the sign includes the wording as a minimum "CAUTION: OXIDIZING GAS(ES) STORED WITHIN NO SMOKING." Storage is planned so cylinders are used in order of which they are received from the supplier. Empty cylinders are segregated from full cylinders. When facility employs cylinders with integral pressure gauge, a threshold pressure</p>			K 923			12/3/21

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K 923	<p>Continued From page 12</p> <p>considered empty is established. Empty cylinders are marked to avoid confusion. Cylinders stored in the open are protected from weather.</p> <p>11.3.1, 11.3.2, 11.3.3, 11.3.4, 11.6.5 (NFPA 99)</p> <p>This REQUIREMENT is not met as evidenced by:</p> <p>Based on observation and staff interview, the facility failed to store medical gas per NFPA 99 (2012 edition), Health Care Facilities Code, sections 11.3.4, 11.6.2.3, 11.6.5 This deficient condition could have a widespread impact on the residents within the facility.</p> <p>Findings include:</p> <p>1. On 09/16/2021 between 11:30 AM to 04:30 PM, it was revealed that RM W-302 - Med Gas storage had mixed storage of cylinders (empty/full) and excess combustible storage within 5-feet of cylinders.</p> <p>2. On 09/16/2021 between 11:30 AM to 04:30 PM, it was revealed that RM C-1 - Med Gas storage had mixed storage of cylinders (empty/full).</p> <p>This deficient condition was verified by the Maintenance Director.</p>	K 923	<p>K923</p> <p>I. Corrective Action taken</p> <p>1. RM W 302 Med gas had mixed storage of cylinders and excess combustible storage w/ 5 feet of cylinders: separated full/empty tanks, installed new labeling and cleaned out any combustibles. 10-9-21.</p> <p>II. To ensure deficient practice does not reoccur <input type="checkbox"/> Maintenance Director will inspect through closure date 12-3-21.</p> <p>III. Monitoring <input type="checkbox"/> Maintenance Director will monitor through 12-3-21</p> <p>IV. Responsible : Maintenance Director</p> <p>V. Date of Completion: 12-3-21</p>		

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E 000	Initial Comments	E 000			
	On 9/13/21 through 9/16/21, a survey for compliance with Appendix Z, Emergency Preparedness Requirements, §483.73(b)(6) was conducted during a standard recertification survey. The facility was IN compliance.				
	The facility is enrolled in ePOC and therefore a signature is not required at the bottom of the first page of the CMS-2567 form. Although no plan of correction is required, it is required that the facility acknowledge receipt of the electronic documents.				
F 000	INITIAL COMMENTS	F 000			
	On 9/13/21 through 9/16/21, a standard recertification survey was conducted at your facility. A complaint investigation was also conducted. Your facility was found to be NOT in compliance with the requirements of 42 CFR 483, Subpart B, Requirements for Long Term Care Facilities.				
	The following complaints were found to be SUBSTANTIATED: H5496016C (MN63320), H5496017C (MN73622), H5496018C (MN73624), H5496019C (MN73625), H5496020C (MN73626), H5496021C (MN73627), and H5496022C (MN73764), however NO deficiencies were cited due to actions implemented by the facility prior to survey.				
	The following complaint was found to be UNSUBSTANTIATED: H5496015C (MN54771).				
	The facility's plan of correction (POC) will serve as your allegation of compliance upon the Departments acceptance. Because you are enrolled in ePOC, your signature is not required				

LABORATORY DIRECTOR'S OR PROVIDER/SUPPLIER REPRESENTATIVE'S SIGNATURE

TITLE

(X6) DATE

Electronically Signed

10/20/2021

Any deficiency statement ending with an asterisk (\*) denotes a deficiency which the institution may be excused from correcting providing it is determined that other safeguards provide sufficient protection to the patients. (See instructions.) Except for nursing homes, the findings stated above are disclosable 90 days following the date of survey whether or not a plan of correction is provided. For nursing homes, the above findings and plans of correction are disclosable 14 days following the date these documents are made available to the facility. If deficiencies are cited, an approved plan of correction is requisite to continued program participation.

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F 000	Continued From page 1 at the bottom of the first page of the CMS-2567 form. Your electronic submission of the POC will be used as verification of compliance.	F 000			
F 578 SS=E	<p>Upon receipt of an acceptable electronic POC, an onsite revisit of your facility may be conducted to validate substantial compliance with the regulations has been attained.</p> <p>Request/Refuse/Dscntnue Trmnt;Formlte Adv Dir CFR(s): 483.10(c)(6)(8)(g)(12)(i)-(v)</p> <p>§483.10(c)(6) The right to request, refuse, and/or discontinue treatment, to participate in or refuse to participate in experimental research, and to formulate an advance directive.</p> <p>§483.10(c)(8) Nothing in this paragraph should be construed as the right of the resident to receive the provision of medical treatment or medical services deemed medically unnecessary or inappropriate.</p> <p>§483.10(g)(12) The facility must comply with the requirements specified in 42 CFR part 489, subpart I (Advance Directives).</p> <p>(i) These requirements include provisions to inform and provide written information to all adult residents concerning the right to accept or refuse medical or surgical treatment and, at the resident's option, formulate an advance directive.</p> <p>(ii) This includes a written description of the facility's policies to implement advance directives and applicable State law.</p> <p>(iii) Facilities are permitted to contract with other entities to furnish this information but are still legally responsible for ensuring that the requirements of this section are met.</p> <p>(iv) If an adult individual is incapacitated at the</p>	F 578		10/27/21	

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NAME OF PROVIDER OR SUPPLIER  <b>MINNEOTA MANOR HEALTH CARE CENTER</b>			STREET ADDRESS, CITY, STATE, ZIP CODE <b>700 NORTH MONROE STREET</b> <b>MINNEOTA, MN 56264</b>		
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F 578	<p>Continued From page 2</p> <p>time of admission and is unable to receive information or articulate whether or not he or she has executed an advance directive, the facility may give advance directive information to the individual's resident representative in accordance with State Law.</p> <p>(v) The facility is not relieved of its obligation to provide this information to the individual once he or she is able to receive such information. Follow-up procedures must be in place to provide the information to the individual directly at the appropriate time.</p> <p>This REQUIREMENT is not met as evidenced by:</p> <p>Based on interview and document review the facility failed to ensure appropriately signed physician orders for advanced directives (AD) for 5 of 23 residents (R1, R5, R8, R9, and R20).</p> <p>Findings include:</p> <p>R1's 5/26/21, quarterly Minimum Data Set (MDS) assessment identified R1's cognition was intact.</p> <p>R1's 7/6/21, physician orders failed to include advance directive (AD) orders or physician ordered life-sustaining treatment (POLST).</p> <p>R1's current, undated care plan identified her code status as Full Code.</p> <p>R1's 8/21/20, Cardio Pulmonary Resuscitation (CPR) Consent form was signed by R1 and witnessed by facility registered nurse (RN), but did not contain a physician (MD) signature. Current MD orders also contained no information on AD/CPR status.</p> <p>R5's 6/28/21, annual MDS identified R5 had</p>	F 578	<p>F578</p> <p>1. R1 <input type="checkbox"/> request for signed physician orders for advanced directives faxed 10-26-21</p> <p>R5 - request for signed physician orders for advanced directives faxed 10-26-21</p> <p>R8 <input type="checkbox"/> Resident discharged from facility 10-21-21</p> <p>R9 - request for signed physician orders for advanced directives faxed 10-26-21</p> <p>R20 - request for signed physician orders for advanced directives faxed 10-26-21</p> <p>2. 11 remaining residents charts will be reviewed for a signed physician order for advanced directive <input type="checkbox"/> any resident that does not have a signed order: request for signed physician orders for advanced directives will be faxed 10-26-21</p> <p>3. At any time the resident requests to change their advanced directive, the Care Coordinator will fax a new request for a signed physician order.</p> <p>4. Care Coordinator will report any resident changes in their advance</p>		



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F 578	<p>Continued From page 3</p> <p>moderate cognitive impairment.</p> <p>R5's physician (MD) progress notes and physician orders identified on 3/22/21, R5's Medicare 60 day physician's note failed to include AD orders or a Physician Ordered Life-Sustaining Treatment (POLST) order. Neither virtual or in-person provider visits after this date contained an order or provider signature authorizing AD.</p> <p>R5's current, undated care plan identified her code status as Do Not Resuscitate (DNR).</p> <p>R5's 7/15/21, Cardio Pulmonary Resuscitation (CPR) consent form identified it contained signatures of 2 family members and was witnessed by 2 RN's. The form failed to contain an MD signature.</p> <p>R8's 7/9/2, quarterly MDS identified R8 had intact cognition.</p> <p>R8's MD progress notes and physician orders identified no signed MD order for AD and/ or any code status.</p> <p>R8's 8/21/12, CPR consent form identified the form was signed by R8 and 2 RN's. There was no signature from the MD identifying they were made aware or agreed with R8's code status of DNR.</p> <p>R9's 7/12/21, quarterly MDS identified his cognition was intact.</p> <p>R9's current, undated care plan identified his resuscitation status as Full Code.</p> <p>R9's 4/6/21 CPR Consent form contained R9's signature, but no witness signatures had been</p>	F 578	<p>directive to the D.O.N. who will monitor and double check that the physician order request has been sent. Weekly 10/26/21 until closure 12/3/2021.</p>		

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F 578	<p>Continued From page 4 included on the document.</p> <p>R9's MD progress notes and physician orders identified on 9/14/21, during a virtual provider visit, there was no mention of the AD and/or CPR status having been ordered and signed by the MD</p> <p>R20's 8/11/21, Significant Change MDS identified R20 had severe cognitive impairment.</p> <p>R20's current, undated care plan identified his resuscitation status as DNR.</p> <p>R20's 6/11/21, CPR Consent form identified No CPR and was signed only by a family member and a staff RN. No MD signature was acquired.</p> <p>R20's MD progress notes and physician orders identified on 8/13/21, during a virtual provider visit, there was no mention of the AD and/or CPR status having been ordered and signed by the MD.</p> <p>Interview on 9/14/21 at 2:35 p.m., with the medical director identified he expected the DNR/DNI and code status for each resident to be included in the list of orders and treatments reviewed and signed by the MD at the time of admission to the facility and renewed with scheduled provider visits. If there was a change in the resident's advance directive choice then he would expect the MD to be notified and signed MD orders obtained as soon as possible.</p> <p>Interview on 9/15/21 at 4:06 p.m. with RN-A identified the facility process for completion of the CPR Consent form was explanation of what CPR, and DNR/DNI meant to the resident, question the resident and/or their representative of code status</p>	F 578			

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F 578	Continued From page 5 preference. Once the choice was made, the CPR consent form was completed, signed by the resident/family member and witnessed by the staff nurse who reviewed the information. The nurse would then enter the resident choice into the electronic medical record under the section for AD. This was completed by checking the appropriate box, which was saved on the system and populated the top of resident documents with the identification information along with the chosen code status. RN-A identified according to facility policy, an MD order was not required as it was felt to be a resident and/or family choice.  Interview on 9/16/21 at 8:30 a.m., the director of nursing (DON) identified the facility did not routinely request signed MD orders for code status. At times the provider would occasionally note the AD and include it in the signed progress note or orders, but it was not consistently included.  A policy related to AD was requested but not provided by the end of the survey.	F 578			
F 656 SS=D	Develop/Implement Comprehensive Care Plan CFR(s): 483.21(b)(1)  §483.21(b) Comprehensive Care Plans §483.21(b)(1) The facility must develop and implement a comprehensive person-centered care plan for each resident, consistent with the resident rights set forth at §483.10(c)(2) and §483.10(c)(3), that includes measurable objectives and timeframes to meet a resident's medical, nursing, and mental and psychosocial needs that are identified in the comprehensive assessment. The comprehensive care plan must describe the following -	F 656			10/27/21

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F 656	<p>Continued From page 6</p> <p>(i) The services that are to be furnished to attain or maintain the resident's highest practicable physical, mental, and psychosocial well-being as required under §483.24, §483.25 or §483.40; and</p> <p>(ii) Any services that would otherwise be required under §483.24, §483.25 or §483.40 but are not provided due to the resident's exercise of rights under §483.10, including the right to refuse treatment under §483.10(c)(6).</p> <p>(iii) Any specialized services or specialized rehabilitative services the nursing facility will provide as a result of PASARR recommendations. If a facility disagrees with the findings of the PASARR, it must indicate its rationale in the resident's medical record.</p> <p>(iv) In consultation with the resident and the resident's representative(s)-</p> <p>(A) The resident's goals for admission and desired outcomes.</p> <p>(B) The resident's preference and potential for future discharge. Facilities must document whether the resident's desire to return to the community was assessed and any referrals to local contact agencies and/or other appropriate entities, for this purpose.</p> <p>(C) Discharge plans in the comprehensive care plan, as appropriate, in accordance with the requirements set forth in paragraph (c) of this section.</p> <p>This REQUIREMENT is not met as evidenced by:</p> <p>Based on interview, and document review the facility failed to ensure a comprehensive care plan was developed for psychoactive medication use that included target behaviors and individualized interventions for 2 of 5 residents (R1 and R17).</p> <p>Findings include:</p>	F 656	<p>F656</p> <p>1. R1 <input type="checkbox"/> Comprehensive care plan will be updated to include target behaviors and individualized interventions. Care plan update 10-26-21. Resident discharging on 10-27-21.</p> <p>R17 <input type="checkbox"/> discharged from facility 10-19-21</p>		

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F 656	<p>Continued From page 7</p> <p>R1's undated, Resident Face Sheet, identified R1 had been admitted to the facility originally in August 2020, with the the most recent re-admission in January 2021. R1 had a diagnosis of depression and anxiety.</p> <p>R1's 5/26/21, quarterly Minimum Data Set (MDS) assessment identified R1's cognition was intact. In the assessment, R1 was identified as feeling bad about themselves, felt they were a failure, or have let themselves or their family down for 2-6 days during the look-back period. R1 had trouble concentrating on things, such as reading the newspaper or watching television for 2-6 day during the look-back period. R1 needed total assistance for transfers, dressing, and locomotion, was always incontinent of bladder and bowel, took insulin, anti-anxiety, anti-depression, and diuretic (fluid pills) medication daily.</p> <p>R1's 7/6/21, physician orders identified R1 was to be administered buspirone (anti-depressant) 15 milligrams (mg) twice a day for anxiety disorder, starting 1/5/21, and paroxetine HCl 20 mg once each evening for anxiety disorder, starting 1/23/21. There were no identified anxiety symptoms R1 was diagnosed with, associated with those medicaion orders.</p> <p>R1's 8/13/21, care plan identified R1 was still unfamiliar with their nursing home routine, and continued to adjust to facility since September 2020. Staff were to allow R1 individual choices in their daily routine, assist R1 in quickly resolving any problems that arose, and gently help R1 be realistic in goal setting. R1 was at risk for injury related to psychotropic medication and was to be</p>	F 656	<p>2. 14 remaining residents care plans will be reviewed to ensure a comprehensive care plan is developed for psychoactive medication use that include target behaviors and individualized interventions. Care Plans will be updated 10-26-21.</p> <p>3. Facility will monitor any changes residents have regarding psychoactive medications and update the care plans to ensure target behaviors and individualized interventions are in place. Care Coordinator will review weekly to ensure the care plan is updated and documentation is complete.</p> <p>4. Weekly report will be reviewed with D.O.N. to ensure compliance. 10/26/21 until closure 12/3/21.</p>		

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F 656	<p>Continued From page 8</p> <p>be free of significant side effects for 90 days. Interventions were discussed with R1 and family. R1's family felt there was benefit to taking their medication and agreed to on-going use and have signed the informed consent. R1 was to take medications as ordered by physician. The pharmacy consult was to achieve the lowest dose necessary and would recommend trial reductions as deemed appropriate. Staff were to watch for and report side effects. Periodically staff were to perform a review of (targeted) behaviors to ensure R1 was receiving the appropriate dose. There was no mention what the target behaviors were, when they would be reviewed, or how the facility would determine if medication was appropriate or therapeutic.</p> <p>Interview on 9/15/21 at 10:09 a.m., with trained medication aide (TMA)-A identified she was unaware of any specific symptoms or behaviors staff were to watch for. She revealed R1 had no behaviors but would tend to "worry a lot".</p> <p>Interview on 9/15/21 at 2:37 p.m., with R1 identified she took anti-anxiety medication as she had a lot of anxiety and a little OCD (obsessive compulsive disorder). She liked things "a certain way". She said she liked to be in control and if that did not happen she would become very anxious. She felt the medication she took helped her with her anxiety.</p> <p>R17's undated, Resident Face Sheet identified R17 had been admitted to the facility in February 2020 with a re-admission in January 2021. R17 had a diagnosis of depression.</p> <p>R17's 8/6/21, quarterly Minimum Data Set (MDS) assessment identified R17's cognition was intact.</p>	F 656			

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F 656	<p>Continued From page 9</p> <p>R17 had trouble falling asleep, staying asleep, and/or sleeping too much. R17 had poor appetite or would overeat 2-6 days during the look-back period. R17 took anti-depressant medication.</p> <p>R17's 5/17/21, care plan identified staff were to allow resident individual choices in daily routine and where she wanted to spend her day. Staff were assist R17 to quickly resolve any problems, and encourage involvement with other residents and activities. Staff were to gently assist R17 be realistic in goal-setting. R17 was at risk for injury related to psychotropic medication and was to be free of significant side effects for 90 days. Interventions were discussed with R17 and family. R17's family felt there was benefit to taking their medication and agreed to on-going use and have signed the informed consent. R1 was to take medications as ordered by the physician. The pharmacy consult was to achieve the lowest dose necessary and would recommend trial reductions as deemed appropriate. Staff were to watch for and report side effects. Periodically staff were to perform a review of (targeted) behaviors to ensure R17 was receiving the appropriate dose. There was no mention what the target behaviors were, when they would be reviewed, or how the facility would determine if medication was appropriate or therapeutic.</p> <p>R17's 7/30/21, Physician Order Report identified citalopram 5 mg once a day for major depressive disorder, which was started on 6/11/21. There were no identified depressive symptoms R1 was diagnosed with, associated with those medicaion orders.</p> <p>Interview on 9/15/21 at 9:10 a.m., with R17</p>	F 656			

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F 656	<p>Continued From page 10</p> <p>identified she had been feeling down and depressed when she first moved to the facility but that had gotten better so she felt the medicaion "must be helping".</p> <p>Interview on 9/15/21 at 10:10 a.m., with TMA-A identified R17 was independent and she was unaware of any symptoms staff were to watch for and report related to her depression. She stated R17 loved to visit and liked to go to activities.</p> <p>Interview on 9/16/21 at 9:35 a.m., with registered nurse (RN)-B identified residents who receive psychoactive medication were to have targeted behaviors identified on the medication administration record (MAR). The nurse would document any behaviors in the residents progress notes. RN-B confirmed neither R1 nor R17 had identified target behaviors listed on the MAR or included on their care plans. RN-B agreed it would be hard to determine if a psychotropic medication would be therapeutic if there were no identified target behaviors to monitor for improvement.</p> <p>Interview on 9/16/21 at 2:45 p.m., with the director of nursing (DON) identified the facility should be identifying target behaviors on the care plan for any resident receiving a psychotropic medication. She further, identified that the medication should be monitored for side effects. She revealed that the target behaviors should be charted on to know if there had been improvement or not with use of the medication. The DON confirmed her expectation would be a resident receiving a psychotropic medication would have identified target behaviors for the medication use and an individualized care plan with interventions for those target behaviors.</p>	F 656			



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F 656	Continued From page 11  Review of 9/16/21, Psychotropic Medication on Admit policy identified the facility was to review all anti-anxiety, hypnotic, and anti-psychotic medication with the resident's physician and work to achieve the lowest dose needed for the resident's well-being. The registered nurse case manager would obtain information about the medication and why it was started. The facility would then communicate with the provider about trial reductions and/or on-going use of the medication. The facility further, would monitor for side effects of medication.	F 656			

<b>FIRE SAFETY SURVEY REPORT - 2012 LIFE SAFETY CODE HEALTHCARE</b>	1. (A) PROVIDER NUMBER  K1	1. (B) MEDICAID I.D. NO.  K2
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PART I — Life Safety Code, New and Existing  
PART II — Health Care Facilities Code, New and Existing  
PART III — Recommendation for Waiver  
PART IV — Crucial Data Extract

OPTIONAL — Chapter 4 – NFPA 101A - Fire Safety Evaluation System for Health Care Occupancies – CMS-2786T

Identifying information as shown in applicable records. Enter changes, if any, alongside each item, giving date of change.

2. NAME OF FACILITY	2. (A) MULTIPLE CONSTRUCTION (BLDGS) A. BUILDING _____ B. WING _____ C. FLOOR _____  K3	2. (B) ADDRESS OF FACILITY (STREET, CITY, STATE, ZIP CODE)	A. <input type="checkbox"/> Fully Sprinklered (All required areas are sprinklered) B. <input type="checkbox"/> Partially Sprinklered (Not all required areas are sprinklered) C. <input type="checkbox"/> None (No sprinkler system) K0180
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3. SURVEY FOR  <input type="checkbox"/> MEDICARE <input type="checkbox"/> MEDICAID	4. DATE OF SURVEY  K4	DATE OF PLAN APPROVAL  K6	SURVEY UNDER 5. <input type="checkbox"/> 2012 EXISTING      6. <input type="checkbox"/> 2012 NEW K7
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5. SURVEY FOR CERTIFICATION OF

1. ☐ HOSPITAL      2. ☐ SKILLED/NURSING FACILITY      4. ☐ ICF/IID UNDER HEALTH CARE      5. ☐ HOSPICE

IF "2" OR "5" ABOVE IS MARKED, CHECK APPROPRIATE ITEM(S) BELOW

1. ☐ ENTIRE FACILITY    2. ☐ DISTINCT PART OF (SPECIFY) \_\_\_\_\_

3. ☐ IF DISTINCT PART OF HOSPITAL, IS HOSPITAL ACCREDITED?



- a. ☐ YES      b. ☐ NO

6. BED COMPOSITION a. TOTAL NO. OF BEDS IN THE FACILITY _____	b. NUMBER OF HOSPITAL BEDS CERTIFIED FOR MEDICARE _____	c. NUMBER OF SKILLED BEDS CERTIFIED FOR MEDICARE _____	d. NUMBER OF SKILLED BEDS CERTIFIED FOR MEDICAID _____	e. NUMBER OF NF or ICF/IID BEDS CERTIFIED FOR MEDICAID _____
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7. A. ☐ THE FACILITY MEETS THE STANDARD, BASED UPON (CHECK ALL APPROPRIATE BOXES)

1. ☐ COMPLIANCE WITH ALL PROVISIONS    2. ☐ ACCEPTANCE OF A PLAN OF CORRECTION    3. ☐ RECOMMENDED WAIVERS    4. ☐ FSES    5. ☐ PERFORMANCE BASED DESIGN

B. ☐ THE FACILITY DOES NOT MEET THE STANDARD

SURVEYOR (Signature) 	TITLE	OFFICE	DATE
SURVEYOR ID K10			
FIRE AUTHORITY OFFICIAL  37009	TITLE	OFFICE	DATE

CMS FORMS SHALL BE COMPLETED AND RETAINED AS PART OF THE SURVEY RECORD.

ID PREFIX		MET	NOT MET	N/A	REMARKS
	<b>PART I – NFPA 101 LSC REQUIREMENTS</b> (Items in italics relate to the FSES)				
	<b>SECTION 1 – GENERAL REQUIREMENTS</b>				
K100	<b>General Requirements – Other</b> List in the REMARKS section any LSC Section 18.1 and 19.1 General Requirements that are not addressed by the provided K-tags, but are deficient. This information, along with the applicable Life Safety Code or NFPA standard citation, should be included on Form CMS-2567.				
K111	<b>Building Rehabilitation</b> <i>Repair, Renovation, Modification, or Reconstruction</i> Any building undergoing repair, renovation, modification, or reconstruction complies with both of the following: <ul style="list-style-type: none"> <li>• Requirements of Chapter 18 and 19.</li> <li>• Requirements of the applicable Sections 43.3, 43.4, 43.5, and 43.6.</li> </ul> 18.1.1.4.3, 19.1.1.4.3, 43.1.2.1 <b>Change of Use or Change of Occupancy</b> Any building undergoing change of use or change of occupancy classification complies with the requirements of Section 43.7, unless permitted by 18.1.1.4.2 or 19.1.1.4.2. 18.1.1.4.2 (4.6.7 and 4.6.11), 19.1.1.4.2 (4.6.7 and 4.6.11), 43.1.2.2 (43.7) <b>Additions</b> Any building undergoing an addition shall comply with the requirements of Section 43.8. If the building has a common wall with a nonconforming building, the common wall is a fire barrier having at least a two hour fire resistance rating constructed of materials as required for the addition. Communicating openings occur only in corridors and are protected by approved self-closing fire doors with at least a 1-1/2 hour fire resistance rating. Additions comply with the requirements of Section 43.8. 18.1.1.4.1 (4.6.7 and 4.6.11), 18.1.1.4.1.1 (8.3), 18.1.1.4.1.2, 18.1.1.4.1.3, 19.1.1.4.1 (4.6.7 and 4.6.11), 19.1.1.4.1.1 (8.3), 19.1.1.4.1.2, 19.1.1.4.1.3, 43.1.2.3(43.8)				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K112	<b>Sprinkler Requirements for Major Rehabilitation</b> If a nonsprinklered smoke compartment has undergone major rehabilitation the automatic sprinkler requirements of 18.3.5 have been applied to the smoke compartment. In cases where the building is not protected throughout by a sprinkler system, the requirements of 18.4.3.2, 18.4.3.3, and 18.4.3.8 are also met. Note: Major rehabilitation involves the modification of more than 50 percent, or more than 4500 ft <sup>2</sup> of the area of the smoke compartment. 18.1.1.4.3.3, 19.1.1.4.3.3				
K131	<b>Multiple Occupancies – Sections of Health Care Facilities</b> Sections of health care facilities classified as other occupancies meet all of the following: <ul style="list-style-type: none"> <li>• They are not intended to serve four or more inpatients for purposes of housing, treatment, or customary access.</li> <li>• They are separated from areas of health care occupancies by construction having a minimum two hour fire resistance rating in accordance with Chapter 8.</li> <li>• The entire building is protected throughout by an approved, supervised automatic sprinkler system in accordance with Section 9.7.</li> </ul> Hospital outpatient surgical departments are required to be classified as an Ambulatory Health Care Occupancy regardless of the number of patients served. 18.1.3.3, 19.1.3.3, 42 CFR 482.41, 42 CFR 485.623				
K132	<b>Multiple Occupancies – Contiguous Non-Health Care Occupancies</b> Non-health care occupancies that are located immediately next to a Health Care Occupancy, but are primarily intended to provide outpatient services are permitted to be classified as Business or Ambulatory Health Care Occupancies, provided the facilities are separated by construction having not less than two hour fire resistance-rated construction, and are not intended to provide services simultaneously for four or more inpatients. Outpatient surgical departments must be classified as Ambulatory Health Care Occupancy regardless of the number of patients served. 18.1.3.4.1, 19.1.3.4.1				

ID PREFIX		MET	NOT MET	N/A	REMARKS																																
K133	<p><b>Multiple Occupancies – Construction Type</b></p> <p>Where separated occupancies are in accordance with 18/19.1.3.2 or 18/19.1.3.4, the most stringent construction type is provided throughout the building, unless a two hour separation is provided in accordance with 8.2.1.3, in which case the construction type is determined as follows:</p> <ul style="list-style-type: none"> <li>The construction type and supporting construction of the health care occupancy is based on the story in which it is located in the building in accordance with 18/19.1.6 and Tables 18/19.1.6.1.</li> <li>The construction type of the areas of the building enclosing the other occupancies shall be based on the applicable occupancy chapters.</li> </ul> <p>18.1.3.5, 19.1.3.5, 8.2.1.3</p>																																				
K161	<p><b>Building Construction Type and Height</b></p> <p>2012 EXISTING</p> <p>Building construction type and stories meets Table 19.1.6.1, unless otherwise permitted by 19.1.6.2 through 19.1.6.7</p> <p>19.1.6.4, 19.1.6.5</p> <table border="1"> <thead> <tr> <th></th> <th></th> <th>Construction Type</th> <th></th> </tr> </thead> <tbody> <tr> <td>1</td> <td></td> <td>I (442), I (332), II (222)</td> <td>Any number of stories non-sprinklered or sprinklered</td> </tr> <tr> <td>2</td> <td></td> <td>II (111)</td> <td>One story non-sprinklered Maximum 3 stories sprinklered</td> </tr> <tr> <td>3</td> <td></td> <td>II (000)</td> <td rowspan="4">Not allowed non-sprinklered Maximum 2 stories sprinklered</td> </tr> <tr> <td>4</td> <td></td> <td>III (211)</td> </tr> <tr> <td>5</td> <td></td> <td>IV (2HH)</td> </tr> <tr> <td>6</td> <td></td> <td>V (111)</td> </tr> <tr> <td>7</td> <td></td> <td>III (200)</td> <td rowspan="2">Not allowed non-sprinklered Maximum 1 story sprinklered</td> </tr> <tr> <td>8</td> <td></td> <td>V (000)</td> </tr> </tbody> </table> <p><i>Sprinklered stories must be sprinklered throughout by an approved, supervised automatic system in accordance with section 9.7. (See 19.3.5)</i></p> <p><i>Give a brief description, in REMARKS, of the construction, the number of stories, including basements, floors on which patients are located, location of smoke or fire barriers and dates of approval. Complete sketch or attach small floor plan of the building as appropriate.</i></p>			Construction Type		1		I (442), I (332), II (222)	Any number of stories non-sprinklered or sprinklered	2		II (111)	One story non-sprinklered Maximum 3 stories sprinklered	3		II (000)	Not allowed non-sprinklered Maximum 2 stories sprinklered	4		III (211)	5		IV (2HH)	6		V (111)	7		III (200)	Not allowed non-sprinklered Maximum 1 story sprinklered	8		V (000)				
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ID PREFIX		MET	NOT MET	N/A	REMARKS																							
K161	<p>2012 NEW</p> <p>Building construction type and stories meets Table 18.1.6.1, unless otherwise permitted by 18.1.6.2 through 18.1.6.7</p> <p>18.1.6.4, 18.1.6.5</p> <table border="1"> <thead> <tr> <th></th> <th>Construction Type</th> <th></th> </tr> </thead> <tbody> <tr> <td>1</td> <td>I (442), I (332), II (222)</td> <td>Not allowed non-sprinklered Any number of stories sprinklered</td> </tr> <tr> <td>2</td> <td>II (111)</td> <td>Not allowed non-sprinklered Maximum 3 stories sprinklered</td> </tr> <tr> <td>3</td> <td>II (000)</td> <td rowspan="4">Not allowed non-sprinklered Maximum 1 story sprinklered</td> </tr> <tr> <td>4</td> <td>III (211)</td> </tr> <tr> <td>5</td> <td>IV (2HH)</td> </tr> <tr> <td>6</td> <td>V (111)</td> </tr> <tr> <td>7</td> <td>III (200)</td> <td rowspan="2">Not allowed non-sprinklered</td> </tr> <tr> <td>8</td> <td>V (000)</td> </tr> </tbody> </table> <p><i>Sprinklered stories must be sprinklered throughout by an approved, supervised automatic system in accordance with section 9.7. (See 18.3.5)</i></p> <p><i>Give a brief description, in REMARKS, of the construction, the number of stories, including basements, floors on which patients are located, location of smoke or fire barriers and dates of approval. Complete sketch or attach small floor plan of the building as appropriate.</i></p>		Construction Type		1	I (442), I (332), II (222)	Not allowed non-sprinklered Any number of stories sprinklered	2	II (111)	Not allowed non-sprinklered Maximum 3 stories sprinklered	3	II (000)	Not allowed non-sprinklered Maximum 1 story sprinklered	4	III (211)	5	IV (2HH)	6	V (111)	7	III (200)	Not allowed non-sprinklered	8	V (000)				
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K162	<p><b>Roofing Systems Involving Combustibles</b></p> <p>2012 EXISTING</p> <p>Buildings of Type I (442), Type I (332), Type II (222), or Type II (111) having roof systems employing combustible roofing supports, decking or roofing meet the following:</p> <ol style="list-style-type: none"> <li>1. roof covering meets Class C requirements.</li> <li>2. roof is separated from occupied building portions with a noncombustible floor assembly using not less than 2½ inches concrete or gypsum fill.</li> <li>3. attic or other space is either unoccupied or protected throughout by an approved automatic sprinkler system.</li> </ol> <p>19.1.6.2*, ASTM E108, ANSI/UL 790</p>																											

ID PREFIX		MET	NOT MET	N/A	REMARKS
K162	<b>2012 NEW</b> Buildings of Type I (442), Type I (332), Type II (222), Type II (111) having roof systems employing combustible roofing supports, decking or roofing meet the following: 1. roof covering meets Class A requirements. 2. roof is separated from occupied building portions with 2 hour fire resistive noncombustible floor assembly using not less than 2½ inches concrete or gypsum fill. 3. the structural elements supporting the rated floor assembly meet the required fire resistance rating of the building. 18.1.6.2, ASTM E108, ANSI/UL 790				
K163	<b>Interior Nonbearing Wall Construction</b> Interior nonbearing walls in Type I or II construction are constructed of noncombustible or limited-combustible materials. Interior nonbearing walls required to have a minimum 2 hour fire resistance rating are permitted to be fire-retardant-treated wood enclosed within noncombustible or limited-combustible materials, provided they are not used as shaft enclosures. 18.1.6.4, 18.1.6.5, 19.1.6.4, 19.1.6.5				
<b>SECTION 2 – MEANS OF EGRESS REQUIREMENTS</b>					
K200	<b>Means of Egress Requirements – Other</b> List in the REMARKS section any LSC Section 18.2 and 19.2 Means of Egress requirements that are not addressed by the provided K-tags, but are deficient. This information, along with the applicable Life Safety Code or NFPA standard citation, should be included on Form CMS-2567. 18.2, 19.2				
K211	<b>Means of Egress – General</b> Aisles, passageways, corridors, exit discharges, exit locations, and accesses are in accordance with Chapter 7, and the means of egress is continuously maintained free of all obstructions to full use in case of emergency, unless modified by 18/19.2.2 through 18/19.2.11. 18.2.1, 19.2.1, 7.1.10.1				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K221	<b>Patient Sleeping Room Doors</b> Locks on patient sleeping room doors are not permitted unless the key-locking device that restricts access from the corridor does not restrict egress from the patient room, or the locking arrangement is permitted for patient clinical, security or safety needs in accordance with 18.2.2.2.5 or 19.2.2.2.5. 18.2.2.2, 19.2.2.2, TIA 12-4				
K222	<b>Egress Doors</b> Doors in a required means of egress shall not be equipped with a latch or a lock that requires the use of a tool or key from the egress side unless using one of the following special locking arrangements:  <input type="checkbox"/> CLINICAL NEEDS OR SECURITY THREAT LOCKING Where special locking arrangements for the clinical security needs of the patient are used, only one locking device shall be permitted on each door and provisions shall be made for the rapid removal of occupants by: remote control of locks; keying of all locks or keys carried by staff at all times; or other such reliable means available to the staff at all times. 18.2.2.2.5.1, 18.2.2.2.6, 19.2.2.2.5.1, 19.2.2.2.6  <input type="checkbox"/> SPECIAL NEEDS LOCKING ARRANGEMENTS Where special locking arrangements for the safety needs of the patient are used, all of the Clinical or Security Locking requirements are being met. In addition, the locks must be electrical locks that fail safely so as to release upon loss of power to the device; the building is protected by a supervised automatic sprinkler system and the locked space is protected by a complete smoke detection system (or is constantly monitored at an attended location within the locked space); and both the sprinkler and detection systems are arranged to unlock the doors upon activation. 18.2.2.2.5.2, 19.2.2.2.5.2, TIA 12-4				



ID PREFIX		MET	NOT MET	N/A	REMARKS
K222	<p><input type="checkbox"/> DELAYED-EGRESS LOCKING ARRANGEMENTS</p> <p>Approved, listed delayed-egress locking systems installed in accordance with 7.2.1.6.1 shall be permitted on door assemblies serving low and ordinary hazard contents in buildings protected throughout by an approved, supervised automatic fire detection system or an approved, supervised automatic sprinkler system.</p> <p>18.2.2.2.4, 19.2.2.2.4</p> <p><input type="checkbox"/> ACCESS-CONTROLLED EGRESS LOCKING ARRANGEMENTS</p> <p>Access-Controlled Egress Door assemblies installed in accordance with 7.2.1.6.2 shall be permitted.</p> <p>18.2.2.2.4, 19.2.2.2.4</p> <p><input type="checkbox"/> ELEVATOR LOBBY EXIT ACCESS LOCKING ARRANGEMENTS</p> <p>Elevator lobby exit access door locking in accordance with 7.2.1.6.3 shall be permitted on door assemblies in buildings protected throughout by an approved, supervised automatic fire detection system and an approved, supervised automatic sprinkler system.</p> <p>18.2.2.2.4, 19.2.2.2.4</p>				
K223	<p><b>Doors with Self-Closing Devices</b></p> <p>Doors in an exit passageway, stairway enclosure, or horizontal exit, smoke barrier, or hazardous area enclosure are self-closing and kept in the closed position, unless held open by a release device complying with 7.2.1.8.2 that automatically closes all such doors throughout the smoke compartment or entire facility upon activation of:</p> <ul style="list-style-type: none"> <li>• Required manual fire alarm system; and</li> <li>• Local smoke detectors designed to detect smoke passing through the opening or a required smoke detection system; and</li> <li>• Automatic sprinkler system, if installed; and</li> <li>• Loss of power.</li> </ul> <p>18.2.2.2.7, 18.2.2.2.8, 19.2.2.2.7, 19.2.2.2.8</p>				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K224	<b>Horizontal-Sliding Doors</b> Horizontal-sliding doors permitted by 7.2.1.14 that are not automatic-closing are limited to a single leaf and shall have a latch or other mechanism to ensure the door will not rebound. Horizontal-sliding doors serving an occupant load fewer than 10 shall be permitted, providing all of the following criteria are met: <ul style="list-style-type: none"> <li>• Area served by the door has no high hazard contents.</li> <li>• Door is operable from either side without special knowledge or effort.</li> <li>• Force required to operate the door in the direction of travel is <math>\leq 30</math> lbf to set the door in motion and <math>\leq 15</math> lbf to close or open to the required width.</li> <li>• Assembly is appropriately fire rated, and where rated, is self-or automatic-closing by smoke detection per 7.2.1.8, and installed per NFPA 80.</li> <li>• Where required to latch, the door has a latch or other mechanism to ensure the door will not rebound.</li> </ul> 18.2.2.2.10, 19.2.2.2.10				
K225	<b>Stairways and Smokeproof Enclosures</b> Stairways and Smokeproof enclosures used as exits are in accordance with 7.2. 18.2.2.3, 18.2.2.4, 19.2.2.3, 19.2.2.4, 7.2				
K226	<b>Horizontal Exits</b> Horizontal exits, if used, are in accordance with 7.2.4 and the provisions of 18.2.2.5.1 through 18.2.2.5.7, or 19.2.2.5.1 through 19.2.2.5.4. 18.2.2.5, 19.2.2.5				
K227	<b>Ramps and Other Exits</b> Ramps, exit passageways, fire and slide escapes, alternating tread devices, and areas of refuge are in accordance with the provisions 7.2.5 through 7.2.12. 18.2.2.6 to 18.2.2.10 or 19.2.2.6 to 19.2.2.10				
K231	<b>Means of Egress Capacity</b> The capacity of required means of egress is in accordance with 7.3. 18.2.3.1, 19.2.3.1				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K232	<b>Aisle, Corridor or Ramp Width</b> 2012 EXISTING The width of aisles or corridors (clear or unobstructed) serving as exit access shall be at least 4 feet and maintained to provide the convenient removal of nonambulatory patients on stretchers, except as modified by 19.2.3.4, exceptions 1-5. 19.2.3.4, 19.2.3.5				
	2012 NEW The width of aisles or corridors (clear and unobstructed) serving as exit access in hospitals and nursing homes shall be at least 8 feet. In limited care facility and psychiatric hospitals, width of aisles or corridors shall be at least 6 feet, except as modified by the 18.2.3.4 or 18.2.3.5 exceptions. 18.2.3.4, 18.2.3.5				
K233	<b>Clear Width of Exit and Exit Access Doors</b> 2012 EXISTING Exit access doors and exit doors are of the swinging type and are at least 32 inches in clear width. Exceptions are provided for existing 34-inch doors and for existing 28-inch doors where the fire plan does not require evacuation by bed, gurney, or wheelchair. 19.2.3.6, 19.2.3.7				
	2012 NEW Exit access doors and exit doors are of the swinging type and are at least 41.5 inches in clear width. In psychiatric hospitals or limited care facilities, doors are at least 32 inches wide. Doors not subject to patient use, in exit stairway enclosures, or serving newborn nurseries shall be no less than 32 inches in clear width. If using a pair of doors, the doors shall be provided with a rabbet, bevel, or astragal at the meeting edge, at least one of the doors shall provide 32 inches in clear width, and the inactive leaf of the pair shall be secured with automatic flush bolts. 18.2.3.6, 18.2.3.7				
K241	<b>Number of Exits – Story and Compartment</b> Not less than two exits, remote from each other, and accessible from every part of every story are provided for each story. Each smoke compartment shall likewise be provided with two distinct egress paths to exits that do not require the entry into the same adjacent smoke compartment. 18.2.4.1-18.2.4.4, 19.2.4.1-19.2.4.4				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K251	<b>Dead-End Corridors and Common Path of Travel</b> 2012 EXISTING Dead-end corridors shall not exceed 30 feet. Existing dead-end corridors greater than 30 feet shall be permitted to be continued to be used if it is impractical and unfeasible to alter them. 19.2.5.2				
K251	2012 NEW Dead-end corridors shall not exceed 30 feet. Common path of travel shall not exceed 100 feet. 18.2.5.2, 18.2.5.3				
K252	<b>Number of Exits – Corridors</b> Every corridor shall provide access to not less than two approved exits in accordance with Sections 7.4 and 7.5 without passing through any intervening rooms or spaces other than corridors or lobbies. 18.2.5.4, 19.2.5.4				
K253	<b>Number of Exits – Patient Sleeping and Non-Sleeping Rooms</b> Patient sleeping rooms of more than 1,000 square feet or nonsleeping rooms of more than 2,500 square feet have at least two exit access doors remotely located from each other. 18.2.5.5.1, 18.2.5.5.2, 19.2.5.5.1, 19.2.5.5.2				
K254	<b>Corridor Access</b> All habitable rooms not within suites have a door leading directly outside to grade or have a door leading to an exit access corridor. Patient sleeping rooms with less than eight patient beds may have one room intervening to reach an exit access corridor provided the intervening room is equipped with an approved automatic smoke detection system. 18.2.5.6.1 through 18.2.5.6.4, 19.2.5.6.1 through 19.2.5.6.4				
K255	<b>Suite Separation, Hazardous Content, and Subdivision</b> All suites are separated from the remainder of the building (including from other suites) by construction meeting the separation provisions for corridor construction (18.3.6.2-18.3.6.5 or 19.3.6.2-19.3.6.5). Existing approved barriers shall be allowed to continue to be used provided they limit the transfer of smoke. Intervening rooms have no hazardous areas and hazardous areas within suites comply with 18/19.2.5.7.1.3. Subdivision of suites shall be by noncombustible or limited-combustible construction. 18.2.5.7.1.2 through 18.2.5.7.1.4, 19.2.5.7.1.2, 19.2.5.7.1.3, 19.2.5.7.1.4				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K256	<p><b>Sleeping Suites</b></p> <p>Occupants shall have exit access to a corridor or direct access to a horizontal exit. Where <math>\geq 2</math> exits are required, one exit access door may be to a stairway, passageway or to the exterior. Suites shall be provided with constant staff supervision. Staff shall have direct visual supervision of patient sleeping rooms, from a constantly attended location or the room shall be provided with an automatic smoke detection system.</p> <p>Suites more than 1,000 ft<sup>2</sup> shall have 2 or more remote exits. One means of egress from the suite shall be to a corridor and one may be into an adjacent suite separated in accordance with corridor requirements.</p> <p>Suites shall not exceed the following size limitations:</p> <ul style="list-style-type: none"> <li>• 5,000 square feet if the suite is not fully smoke detected or fully sprinklered.</li> <li>• 7,500 square feet if the suite is either fully smoke detected or fully sprinklered.</li> <li>• 10,000 square feet if the suite is both fully smoke detected and fully sprinklered and the sleeping rooms have direct supervision from a constantly attended location.</li> </ul> <p>Travel distance between any point in a suite to exit access shall not exceed 100 feet and distance to an exit shall not exceed 150 feet (200 feet if building is fully sprinklered).</p> <p>18.2.5.7.2, 19.2.5.7.2</p>				
K257	<p><b>Non-Sleeping Suites</b></p> <p>Occupants shall have exit access to a corridor or direct access to a horizontal exit. Where <math>\geq 2</math> exits are required, one exit access door may be to a stairway, passageway or to the exterior.</p> <p>Suites more than 2,500 ft<sup>2</sup> shall have 2 or more remote exits. One means of egress from the suite shall be to a corridor and one may be into an adjacent suite separated in accordance with corridor requirements.</p> <p>Suites shall not exceed 10,000 ft<sup>2</sup>.</p> <p>Travel distance between any point in a suite to exit access shall not exceed 100 feet and distance to an exit shall not exceed 150 feet (200 feet if building is fully sprinklered).</p> <p>18.2.5.7.3, 19.2.5.7.3</p>				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K261	<b>Travel Distance to Exits</b> Travel distance (excluding suites) to exits are measured in accordance with 7.6. <ul style="list-style-type: none"> <li>From any point in the room or suite to exit less than or equal to 150 feet (less than or equal to 200 feet if the building is fully sprinklered).</li> <li>Point in a room to room door less than or equal to 50 feet.</li> </ul> 18.2.6, 19.2.6				
K271	<b>Discharge from Exits</b> Exit discharge is arranged in accordance with 7.7, provides a level walking surface meeting the provisions of 7.1.7 with respect to changes in elevation and shall be maintained free of obstructions. Additionally, the exit discharge shall be a hard packed all-weather travel surface. 18.2.7, 19.2.7				
K281	<b>Illumination of Means of Egress</b> Illumination of means of egress, including exit discharge, is arranged in accordance with 7.8 and shall be either continuously in operation or capable of automatic operation without manual intervention. 18.2.8, 19.2.8				
K291	<b>Emergency Lighting</b> Emergency lighting of at least 1-1/2 hour duration is provided automatically in accordance with 7.9. 18.2.9.1, 19.2.9.1				
K292	<b>Life Support Means of Egress</b> 2012 NEW (INDICATE N/A FOR EXISTING) Buildings equipped with or requiring the use of life support systems (electro-mechanical or inhalation anesthetics) have illumination of means of egress, emergency lighting equipment, exit, and directional signs supplied by the life safety branch of the electrical system described in NFPA 99. (Indicate N/A if life support equipment is for emergency purposes only.) 18.2.9.2, 18.2.10.5				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K293	<b>Exit Signage</b> 2012 EXISTING Exit and directional signs are displayed in accordance with 7.10 with continuous illumination also served by the emergency lighting system. 19.2.10.1 (Indicate N/A in one-story existing occupancies with less than 30 occupants where the line of exit travel is obvious.)				
	2012 NEW Exit and directional signs are displayed in accordance with 7.10 with continuous illumination also served by the emergency lighting system. 18.2.10.1				
<b>SECTION 3 – PROTECTION</b>					
K300	<b>Protection – Other</b> List in the REMARKS section any LSC Section 18.3 and 19.3 Protection requirements that are not addressed by the provided K-tags, but are deficient. This information, along with the applicable Life Safety Code or NFPA standard citation, should be included on Form CMS-2567.				
K311	<b>Vertical Openings – Enclosure</b> 2012 EXISTING Stairways, elevator shafts, light and ventilation shafts, chutes, and other vertical openings between floors are enclosed with construction having a fire resistance rating of at least 1-hour. An atrium may be used in accordance with 8.6. 19.3.1.1 through 19.3.1.6 <i>If all vertical openings are properly enclosed with construction providing at least a 2 hour fire resistance rating, also check this box.</i> <input type="checkbox"/>				
	2012 NEW Stairways, elevator shafts, light and ventilation shafts, chutes, and other vertical openings between floors are enclosed with construction having a fire resistance rating of at least 2 hours connecting four or more stories. (1-hour for single story building and buildings up to three stories in height.) An atrium may be used in accordance with 8.6.7. 18.3.1 through 18.3.1.5				

ID PREFIX		MET	NOT MET	N/A	REMARKS																																
K321	<p><b>Hazardous Areas – Enclosure</b></p> <p>2012 EXISTING</p> <p>Hazardous areas are protected by a fire barrier having 1-hour fire resistance rating (with ¾ hour fire rated doors) or an automatic fire extinguishing system in accordance with 8.7.1 or 19.3.5.9. When the approved automatic fire extinguishing system option is used, the areas shall be separated from other spaces by smoke resisting partitions and doors in accordance with 8.4. Doors shall be self-closing or automatic-closing and permitted to have nonrated or field-applied protective plates that do not exceed 48 inches from the bottom of the door.</p> <p><i>Describe the floor and zone locations of hazardous areas that are deficient in REMARKS.</i></p> <p>19.3.2.1, 19.3.5.9</p> <table border="1"> <thead> <tr> <th>Area</th> <th>Automatic Sprinkler</th> <th>Separation</th> <th>N/A</th> </tr> </thead> <tbody> <tr> <td>a. Boiler and Fuel-Fired Heater Rooms</td> <td></td> <td></td> <td></td> </tr> <tr> <td>b. Laundries (larger than 100 sq. ft.)</td> <td></td> <td></td> <td></td> </tr> <tr> <td>c. Repair, Maintenance, and Paint Shops</td> <td></td> <td></td> <td></td> </tr> <tr> <td>d. Soiled Linen Rooms (exceeding 64 gal.)</td> <td></td> <td></td> <td></td> </tr> <tr> <td>e. Trash Collection Rooms (exceeding 64 gal.)</td> <td></td> <td></td> <td></td> </tr> <tr> <td>f. Combustible Storage Rooms/Spaces (over 50 sq. ft.)</td> <td></td> <td></td> <td></td> </tr> <tr> <td>g. Laboratories (if classified as Severe Hazard - see K322)</td> <td></td> <td></td> <td></td> </tr> </tbody> </table>	Area	Automatic Sprinkler	Separation	N/A	a. Boiler and Fuel-Fired Heater Rooms				b. Laundries (larger than 100 sq. ft.)				c. Repair, Maintenance, and Paint Shops				d. Soiled Linen Rooms (exceeding 64 gal.)				e. Trash Collection Rooms (exceeding 64 gal.)				f. Combustible Storage Rooms/Spaces (over 50 sq. ft.)				g. Laboratories (if classified as Severe Hazard - see K322)							
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K321	<p>2012 NEW</p> <p>Hazardous areas are protected in accordance with 18.3.2.1. The areas shall be enclosed with a 1-hour fire-rated barrier, with a <math>\frac{3}{4}</math> hour fire-rated door without windows (in accordance with 8.7.1.1). Doors shall be self-closing or automatic-closing in accordance with 7.2.1.8. Hazardous areas are protected by a sprinkler system in accordance with 9.7, 18.3.2.1, and 8.4.</p> <p><i>Describe the floor and zone locations of hazardous areas that are deficient in REMARKS.</i></p> <p>18.3.2.1, 7.2.1.8, 8.4, 8.7, 9.7</p> <table border="1"> <thead> <tr> <th>Area</th> <th>Automatic Sprinkler</th> <th>Separation</th> <th>N/A</th> </tr> </thead> <tbody> <tr> <td>a. Boiler and Fuel-Fired Heater Rooms</td> <td></td> <td></td> <td></td> </tr> <tr> <td>b. Laundries (larger than 100 sq. ft.)</td> <td></td> <td></td> <td></td> </tr> <tr> <td>c. Repair, Maintenance, and Paint Shops</td> <td></td> <td></td> <td></td> </tr> <tr> <td>d. Soiled Linen Rooms (exceeding 64 gal.)</td> <td></td> <td></td> <td></td> </tr> <tr> <td>e. Trash Collection Rooms (exceeding 64 gal.)</td> <td></td> <td></td> <td></td> </tr> <tr> <td>f. Combustible Storage Rooms/Spaces (over 50 and less than 100 sq. ft.)</td> <td></td> <td></td> <td></td> </tr> <tr> <td>g. Combustible Storage Rooms/Spaces (over 100 sq. ft.)</td> <td></td> <td></td> <td></td> </tr> <tr> <td>h. Laboratories (if classified as Severe Hazard - see K322)</td> <td></td> <td></td> <td></td> </tr> </tbody> </table>	Area	Automatic Sprinkler	Separation	N/A	a. Boiler and Fuel-Fired Heater Rooms				b. Laundries (larger than 100 sq. ft.)				c. Repair, Maintenance, and Paint Shops				d. Soiled Linen Rooms (exceeding 64 gal.)				e. Trash Collection Rooms (exceeding 64 gal.)				f. Combustible Storage Rooms/Spaces (over 50 and less than 100 sq. ft.)				g. Combustible Storage Rooms/Spaces (over 100 sq. ft.)				h. Laboratories (if classified as Severe Hazard - see K322)							
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ID PREFIX		MET	NOT MET	N/A	REMARKS
K322	<p><b>Laboratories</b></p> <p>Laboratories employing quantities of flammable, combustible, or hazardous materials that are considered a severe hazard are protected by 1-hour fire resistance-rated separation, automatic sprinkler system, and are in accordance with 8.7 and with NFPA 99.</p> <p>Laboratories not considered a severe hazard are protected as hazardous areas (see K321).</p> <p>Laboratories using chemicals are in accordance with NFPA 45, <i>Standard on Fire Protection for Laboratories Using Chemicals</i>.</p> <p>Gas appliances are of appropriate design and installed in accordance with NFPA 54. Shutoff valves are marked to identify material they control.</p> <p>Devices requiring medical grade oxygen from the piped distribution system meet the requirements under 11.4.2.2 (NFPA 99).</p> <p>18.3.2.2, 19.3.2.2, 8.7, 8.7.4.1 (LSC)</p> <p>9.3.1.2, 11.4.3.2, 15.4 (NFPA 99)</p>				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K323	<p><b>Anesthetizing Locations</b></p> <p>Areas designated for administration of general anesthesia (i.e., inhalation anesthetics) are in accordance with 8.7 and NFPA 99.</p> <p>Zone valves are: located immediately outside each life-support, critical care, and anesthetizing location of moderate sedation, deep sedation, or general anesthesia for medical gas or vacuum; readily accessible in an emergency; and arranged so shutting off any one anesthetizing location will not affect others.</p> <p>Area alarm panels are provided to monitor all medical gas, medical-surgical vacuum, and piped WAGD systems. Panels are at locations that provide for surveillance, indicate medical gas pressure decreases of 20 percent and vacuum decreases of 12 inch gauge HgV, and provide visual and audible indication. Alarm sensors are installed either on the source side of individual room zone valve box assemblies or on the patient/use side of each of the individual zone box valve assemblies.</p> <p>The EES critical branch supplies power for task illumination, fixed equipment, select receptacles, and select power circuits, and EES equipment system supplies power to ventilation system.</p> <p>Heating, cooling, and ventilation are in accordance with ASHRAE 170. Medical supply and equipment manufacturer's instructions for use are considered before reducing humidity levels to those allowed by ASHRAE, per S&amp;C 13-58.</p> <p>18.3.2.3, 19.3.2.3 (LSC)</p> <p>5.1.4.8.7, 5.1.4.8.7.2, 5.1.9.3, 5.1.9.3.4, 6.4.2.2.4.2 (NFPA 99)</p>				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K324	<b>Cooking Facilities</b> Cooking equipment is protected in accordance with NFPA 96, <i>Standard for Ventilation Control and Fire Protection of Commercial Cooking Operations</i> , unless: <ul style="list-style-type: none"> <li>residential cooking equipment (i.e., small appliances such as microwaves, hot plates, toasters) are used for food warming or limited cooking in accordance with 18.3.2.5.2, 19.3.2.5.2.</li> <li>cooking facilities open to the corridor in smoke compartments with 30 or fewer patients comply with the conditions under 18.3.2.5.3, 19.3.2.5.3, or</li> <li>cooking facilities in smoke compartments with 30 or fewer patients comply with conditions under 18.3.2.5.4, 19.3.2.5.4.</li> </ul> Cooking facilities protected according to NFPA 96 per 9.2.3 are not required to be enclosed as hazardous areas, but shall not be open to the corridor. 18.3.2.5.1 through 18.3.2.5.4, 19.3.2.5.1 through 19.3.2.5.5, 9.2.3, TIA 12-2				
K325	<b>Alcohol Based Hand Rub Dispenser (ABHR)</b> ABHRs are protected in accordance with 8.7.3.1, unless all conditions are met: <ul style="list-style-type: none"> <li>Corridor is at least 6 feet wide.</li> <li>Maximum individual dispenser capacity is 0.32 gallons (0.53 gallons in suites) of fluid and 18 ounces of Level 1 aerosols.</li> <li>Dispensers shall have a minimum of four foot horizontal spacing.</li> <li>Not more than an aggregate of 10 gallons of fluid or 1135 ounces of aerosol are used in a single smoke compartment outside a storage cabinet, excluding one individual dispenser per room.</li> <li>Storage in a single smoke compartment greater than 5 gallons complies with NFPA 30.</li> <li>Dispensers are not installed within 1 inch of an ignition source.</li> <li>Dispensers over carpeted floors are in sprinklered smoke compartments.</li> <li>ABHR does not exceed 95 percent alcohol.</li> <li>Operation of the dispenser shall comply with Section 18.3.2.6(11) or 19.3.2.6(11).</li> <li>ABHR is protected against inappropriate access.</li> </ul> 18.3.2.6, 19.3.2.6, 42 CFR Parts 403, 418, 460, 482, 483, and 485				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K331	<b>Interior Wall and Ceiling Finish</b> 2012 EXISTING Interior wall and ceiling finishes, including exposed interior surfaces of buildings such as fixed or movable walls, partitions, columns, and have a flame spread rating of Class A or Class B. The reduction in class of interior finish for a sprinkler system as prescribed in 10.2.8.1 is permitted. 10.2, 19.3.3.1, 19.3.3.2 <i>Indicate flame spread rating(s).</i> _____				
	2012 NEW Interior wall and ceiling finishes, including exposed interior surfaces of buildings such as fixed or movable walls, partitions and columns have a flame spread rating of Class A. The reduction in class of interior finish for a sprinkler system as prescribed in 10.2.8.1 is permitted. Individual rooms not exceeding four persons may have a Class A or B finish. Lower half of corridor walls, not exceeding 4 feet in height, may have a Class A or B flame spread rating. 10.2, 18.3.3.1, 18.3.3.2 <i>Indicate flame spread rating(s).</i> _____				
K332	<b>Interior Floor Finish</b> 2012 NEW (Indicate N/A for 2012 EXISTING) Interior finishes shall comply with 10.2. Floor finishes in exit enclosures and exit access corridors and spaces not separated by walls that resist the passage of smoke shall be Class I or II. 18.3.3.3.1, 18.3.3.3.2, 18.3.3.3.3, 10.2, 10.2.7.1, 10.2.7.2				
K341	<b>Fire Alarm System – Installation</b> A fire alarm system is installed with systems and components approved for the purpose in accordance with NFPA 70, <i>National Electric Code</i> , and NFPA 72, <i>National Fire Alarm Code</i> to provide effective warning of fire in any part of the building. In areas not continuously occupied, detection is installed at each fire alarm control unit. In new occupancy, detection is also installed at notification appliance circuit power extenders, and supervising station transmitting equipment. Fire alarm system wiring or other transmission paths are monitored for integrity. 18.3.4.1, 19.3.4.1, 9.6, 9.6.1.8				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K342	<b>Fire Alarm System – Initiation</b> Initiation of the fire alarm system is by manual means and by any required sprinkler system alarm, detection device, or detection system. Manual alarm boxes are provided in the path of egress near each required exit. Manual alarm boxes in patient sleeping areas shall not be required at exits if manual alarm boxes are located at all nurse's stations or other continuously attended staff location, provided alarm boxes are visible, continuously accessible, and 200' travel distance is not exceeded. 18.3.4.2.1, 18.3.4.2.2, 19.3.4.2.1, 19.3.4.2.2, 9.6.2.5				
K343	<b>Fire Alarm – Notification</b> 2012 EXISTING Positive alarm sequence in accordance with 9.6.3.4 are permitted in buildings protected throughout by a sprinkler system. Occupant notification is provided automatically in accordance with 9.6.3 by audible and visual signals. In critical care areas, visual alarms are sufficient. The fire alarm system transmits the alarm automatically to notify emergency forces in the event of a fire. 19.3.4.3, 19.3.4.3.1, 19.3.4.3.2, 9.6.4, 9.7.1.1(1)				
	2012 NEW Positive alarm sequence in accordance with 9.6.3.4 are permitted. Occupant notification is provided automatically in accordance with 9.6.3 by audible and visual signals. In critical care areas, visual alarms are sufficient. The fire alarm system transmits the alarm automatically to notify emergency forces in the event of a fire. Annunciation and annunciation zoning for fire alarm and sprinklers shall be provided by audible and visual indicators and zones shall not be larger than 22,500 square feet per zone. 18.3.4.3 through 18.3.4.3.3, 9.6.4				
K344	<b>Fire Alarm – Control Functions</b> The fire alarm automatically activates required control functions and is provided with an alternative power supply in accordance with NFPA 72. 18.3.4.4, 19.3.4.4, 9.6.1, 9.6.5, NFPA 72				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K345	<b>Fire Alarm System – Testing and Maintenance</b> A fire alarm system is tested and maintained in accordance with an approved program complying with the requirements of NFPA 70, <i>National Electric Code</i> , and NFPA 72, <i>National Fire Alarm and Signaling Code</i> . Records of system acceptance, maintenance and testing are readily available. 9.6.1.3, 9.6.1.5, NFPA 70, NFPA 72				
K346	<b>Fire Alarm – Out of Service</b> Where required fire alarm system is out of services for more than 4 hours in a 24 hour period, the authority having jurisdiction shall be notified, and the building shall be evacuated or an approved fire watch shall be provided for all parties left unprotected by the shutdown until the fire alarm system has been returned to service. 9.6.1.6				
K347	<b>Smoke Detection</b> 2012 EXISTING Smoke detection systems are provided in spaces open to corridors as required by 19.3.6.1. 19.3.4.5.2				
	2012 NEW Smoke detection systems are provided in spaces open to corridors as required by 18.3.6.1 In nursing homes, an automatic smoke detection system is installed in the corridors of all smoke compartments containing resident sleeping rooms, unless the resident sleeping rooms have: <ul style="list-style-type: none"> <li>• smoke detection, or</li> <li>• automatic door closing devices with integral smoke detectors on the room side that provide occupant notification.</li> </ul> Such detectors are electrically interconnected to the fire alarm system. 18.3.4.5.2, 18.3.4.5.3				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K351	<b>Sprinkler System – Installation</b> 2012 EXISTING Nursing homes, and hospitals where required by construction type, are protected throughout by an approved automatic sprinkler system in accordance with NFPA 13, <i>Standard for the Installation of Sprinkler Systems</i> . In Type I and II construction, alternative protection measures are permitted to be substituted for sprinkler protection in specific areas where state or local regulations prohibit sprinklers. In hospitals, sprinklers are not required in clothes closets of patient sleeping rooms where the area of the closet does not exceed 6 ft <sup>2</sup> and sprinkler coverage covers the closet footprint as required by NFPA 13, <i>Standard for Installation of Sprinkler Systems</i> . 19.3.5.1, 19.3.5.2, 19.3.5.3, 19.3.5.4, 19.3.5.5, 19.4.2, 19.3.5.10, 9.7, 9.7.1.1(1)				
	2012 NEW Buildings are to be protected throughout by an approved automatic sprinkler system in accordance with NFPA 13, <i>Standard for the Installation of Sprinkler Systems</i> . In Type I and II construction, alternative protection measures are permitted to be substituted for sprinkler protection in specific areas where State and local regulations prohibit sprinklers. Listed quick-response or listed residential sprinklers are used throughout smoke compartments with patient sleeping rooms. In hospitals, sprinklers are not required in clothes closets of patient sleeping rooms where the area of the closet does not exceed 6 ft <sup>2</sup> and sprinkler coverage covers the closet footprint as required by NFPA 13, <i>Standard for Installation of Sprinkler Systems</i> . 18.3.5.1, 18.3.5.4, 18.3.5.5, 18.3.5.6, 9.7, 9.7.1.1(1), 18.3.5.10				
K352	<b>Sprinkler System – Supervisory Signals</b> Automatic sprinkler system supervisory attachments are installed and monitored for integrity in accordance with NFPA 72, <i>National Fire Alarm and Signaling Code</i> , and provide a signal that sounds and is displayed at a continuously attended location or approved remote facility when sprinkler operation is impaired. 9.7.2.1, NFPA 72				



ID PREFIX		MET	NOT MET	N/A	REMARKS
K353	<b>Sprinkler System – Maintenance and Testing</b> Automatic sprinkler and standpipe systems are inspected, tested, and maintained in accordance with NFPA 25, <i>Standard for the Inspection, Testing, and Maintaining of Water-based Fire Protection Systems</i> . Records of system design, maintenance, inspection and testing are maintained in a secure location and readily available. a) Date sprinkler system last checked. _____ b) Who provided system test. _____ c) Water system supply source. _____ <i>Provide in REMARKS information on coverage for any non-required or partial automatic sprinkler system.</i> 9.7.5, 9.7.7, 9.7.8, and NFPA 25				
K354	<b>Sprinkler System – Out of Service</b> Where the sprinkler system is impaired, the extent and duration of the impairment has been determined, areas or buildings involved are inspected and risks are determined, recommendations are submitted to management or designated representative, and the fire department and other authorities having jurisdiction have been notified. Where the sprinkler system is out of service for more than 10 hours in a 24 hour period, the building or portion of the building affected are evacuated or an approved fire watch is provided until the sprinkler system has been returned to service. 18.3.5.1, 19.3.5.1, 9.7.5, 15.5.2 (NFPA 25)				
K355	<b>Portable Fire Extinguishers</b> Portable fire extinguishers are selected, installed, inspected, and maintained in accordance with NFPA 10, <i>Standard for Portable Fire Extinguishers</i> . 18.3.5.12, 19.3.5.12, NFPA 10				
K361	<b>Corridors – Areas Open to Corridor</b> Spaces (other than patient sleeping rooms, treatment rooms and hazardous areas), waiting areas, nurse's stations, gift shops, and cooking facilities, open to the corridor are in accordance with the criteria under 18.3.6.1 and 19.3.6.1. 18.3.6.1, 19.3.6.1				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K362	<b>Corridors – Construction of Walls</b> 2012 EXISTING Corridors are separated from use areas by walls constructed with at least ½ hour fire resistance rating. In fully sprinklered smoke compartments, partitions are only required to resist the transfer of smoke. In nonsprinklered buildings, walls extend to the underside of the floor or roof deck above the ceiling. Corridor walls may terminate at the underside of ceilings where specifically permitted by Code. Fixed fire window assemblies in corridor walls are in accordance with Section 8.3, but in sprinklered compartments there are no restrictions in area or fire resistance of glass or frames. <i>If the walls have a fire resistance rating, give the rating _____ if the walls terminate at the underside of the ceiling, give brief description in REMARKS, describing the ceiling throughout the floor area.</i> 19.3.6.2, 19.3.6.2.7				
	2012 NEW Corridor walls shall form a barrier to limit the transfer of smoke. Such walls shall be permitted to terminate at the ceiling where the ceiling is constructed to limit the transfer of smoke. No fire resistance rating is required for the corridor walls. 18.3.6.2				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K363	<b>Corridor – Doors</b> 2012 EXISTING Doors protecting corridor openings in other than required enclosures of vertical openings, exits, or hazardous areas resist the passage of smoke and are made of 1¾ inch solid-bonded core wood or other material capable of resisting fire for at least 20 minutes. Doors in fully sprinklered smoke compartments are only required to resist the passage of smoke. Corridor doors and doors to rooms containing flammable or combustible materials have positive latching hardware. Roller latches are prohibited by CMS regulation. These requirements do not apply to auxiliary spaces that do not contain flammable or combustible material. Powered doors complying with 7.2.1.9 are permissible if provided with a device capable of keeping the door closed when a force of 5lbf is applied, whether or not power is applied. Clearance between bottom of door and floor covering is not exceeding 1 inch. There is no impediment to the closing of the doors. Hold open devices that release when the door is pushed or pulled are permitted. Nonrated protective plates of unlimited height are permitted. Dutch doors meeting 19.3.6.3.6 are permitted. Door frames shall be labeled and made of steel or other materials in compliance with 8.3, unless the smoke compartment is sprinklered. Fixed fire window assemblies are allowed per 8.3. In sprinklered compartments there are no restrictions in area or fire resistance of glass or frames in window assemblies. 19.3.6.3, 42 CFR Parts 403, 418, 460, 482, 483, and 485 Show in REMARKS details of doors such as fire protection ratings, automatics closing devices, etc.				
	2012 NEW Doors protecting corridor openings shall be constructed to resist the passage of smoke. Corridor doors and doors to rooms containing flammable or combustible materials have self-latching and positive latching hardware. Roller latches are prohibited by CMS regulation. These requirements do not apply to auxiliary spaces that do not contain flammable or combustible material. Powered doors complying with 7.2.1.9 are permissible if provided with a device capable of keeping the door closed when a force of 5lbf is applied, whether or not power is applied. Clearance between bottom of door and floor covering is not exceeding 1 inch. There is no impediment to the closing of the doors. Hold open devices that release when the door is pushed or pulled are permitted. Nonrated protective plates of unlimited height are permitted. Dutch doors meeting 18.3.6.3.6 are permitted. 18.3.6.3, 42 CFR Parts 403, 418, 460, 482, 483, and 485 Show in REMARKS details of doors such as fire protection ratings, automatic closing devices, etc.				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K364	<p><b>Corridor – Openings</b></p> <p>Transfer grilles are not used in corridor walls or doors. Auxiliary spaces that do not contain flammable or combustible materials are permitted to have louvers or be undercut.</p> <p>In other than smoke compartments containing patient sleeping rooms, miscellaneous openings are permitted in vision panels or doors, provided the openings per room do not exceed 20 in<sup>2</sup> and are at or below half the distance from floor to ceiling. In sprinklered rooms, the openings per room do not exceed 80 in<sup>2</sup>.</p> <p>Vision panels in corridor walls or doors shall be fixed window assemblies in approved frames. (In fully sprinklered smoke compartments, there are no restrictions in the area and fire resistance of glass and frames.)</p> <p>18.3.6.5.1, 19.3.6.5.2, 8.3</p>				
K371	<p><b>Subdivision of Building Spaces – Smoke Compartments</b></p> <p>2012 EXISTING</p> <p>Smoke barriers shall be provided to form at least two smoke compartments on every sleeping floor with a 30 or more patient bed capacity. Size of compartments cannot exceed 22,500 square feet or a 200-foot travel distance from any point in the compartment to a door in the smoke barrier.</p> <p>19.3.7.1, 19.3.7.2</p> <p><i>Detail in REMARKS zone dimensions including length of zones and dead-end corridors.</i></p>				
	<p>2012 NEW</p> <p>Smoke barriers shall be provided to form at least two smoke compartments on every floor used by inpatients for sleeping or treatment, and on every floor with an occupant load of 50 or more persons, regardless of use. Size of compartments cannot exceed 22,500 square feet or a 200-foot travel distance from any point in the compartment to a door in the smoke barrier.</p> <p>Smoke subdivision requirements do not apply to any of the stories or areas described in 18.3.7.2.</p> <p>18.3.7.1, 18.3.7.2</p> <p><i>Detail in REMARKS zone dimensions including length of zones and dead-end corridors.</i></p>				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K372	<b>Subdivision of Building Spaces – Smoke Barrier Construction</b> 2012 EXISTING Smoke barriers shall be constructed to a ½ hour fire resistance rating per 8.5. Smoke barriers shall be permitted to terminate at an atrium wall. Smoke dampers are not required in duct penetrations in fully ducted HVAC systems where an approved sprinkler system is installed for smoke compartments adjacent to the smoke barrier. 19.3.7.3, 8.6.7.1(1) <i>Describe any mechanical smoke control system in REMARKS.</i>				
	2012 NEW Smoke barriers shall be constructed to provide at least a 1-hour fire resistance rating and constructed in accordance with 8.5. Smoke barriers shall be permitted to terminate at an atrium wall. Smoke dampers are not required in duct penetrations of fully ducted HVAC systems. 18.3.7.3, 18.3.7.4, 18.3.7.5, 8.3 <i>Describe any mechanical smoke control system in REMARKS.</i>				
K373	<b>Subdivision of Building Spaces – Accumulation Space</b> Space shall be provided on each side of smoke barriers to adequately accommodate the total number of occupants in adjoining compartments. 18.3.7.5.1, 18.3.7.5.2, 19.3.7.5.1, 19.3.7.5.2				
K374	<b>Subdivision of Building Spaces – Smoke Barrier Doors</b> 2012 EXISTING Doors in smoke barriers are 1¾-inch thick solid bonded wood-core doors or of construction that resists fire for 20 minutes. Nonrated protective plates of unlimited height are permitted. Doors are permitted to have fixed fire window assemblies per 8.5. Doors are self-closing or automatic-closing, do not require latching, and are not required to swing in the direction of egress travel. Door opening provides a minimum clear width of 32 in for swinging or horizontal doors. 19.3.7.6, 19.3.7.8, 19.3.7.9				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K374	<p>2012 NEW</p> <p>Doors in smoke barriers have at least a 20-minute fire protection rating or are at least 1¾-inch thick solid bonded core wood.</p> <p>Required clear widths are provided per 18.3.7.6(4) and (5).</p> <p>Nonrated protective plates of unlimited height are permitted. Horizontal-sliding doors comply with 7.2.1.14. Swinging doors shall be arranged so that each door swings in an opposite direction.</p> <p>Doors shall be self-closing and rabbets, bevels, or astragals are required at the meeting edges. Positive latching is not required.</p> <p>18.3.7.6, 18.3.7.7, 18.3.7.8</p>				
K379	<p><b>Smoke Barrier Door Glazing</b></p> <p>2012 EXISTING</p> <p>Openings in smoke barrier doors shall be fire-rated glazing or wired glass panels in steel frames.</p> <p>19.3.7.6, 19.3.7.6.2, 8.5</p>				
	<p>2012 NEW</p> <p>Windows in smoke barrier doors shall be installed in each cross corridor swinging or horizontal-sliding door protected by fire-rated glazing or by wired glass panels in approved frames.</p> <p>18.3.7.9</p>				
K381	<p><b>Sleeping Room Outside Windows and Doors</b></p> <p>Every patient sleeping room has an outside window or outside door. In new occupancies, sill height does not exceed 36 inches above the floor. Windows in atrium walls are considered outside windows. Newborn nurseries and rooms intended for occupancy less than 24 hours have no outside window or door requirements. Window sills in special nursing care areas (e.g., ICU, CCU, hemodialysis, neonatal) do not exceed 60 inches above the floor.</p> <p>42 CFR 403, 418, 460, 482, 483, and 485</p>				
	<b>SECTION 4 – SPECIAL PROVISIONS</b>				
K400	<p><b>Special Provisions – Other</b></p> <p>List in the REMARKS section any LSC Section 18.4 and 19.4 Special Provisions requirements that are not addressed by the provided K-tags, but are deficient. This information, along with the applicable Life Safety Code or NFPA standard citation, should be included on Form CMS-2567.</p>				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K421	<b>High-Rise Buildings</b> 2012 EXISTING High-rise buildings are protected throughout by an approved, supervised automatic sprinkler system in accordance with Section 9.7 within 12 years of LSC final rule effective date. 19.4.2				
	2012 NEW High-rise buildings comply with section 11.8. 18.4.2				
<b>SECTION 5 – BUILDING SERVICES</b>					
K500	<b>Building Services – Other</b> List in the REMARKS section any LSC Section 18.5 and 19.5 Building Services requirements that are not addressed by the provided K-tags, but are deficient. This information, along with the applicable Life Safety Code or NFPA standard citation, should be included on Form CMS-2567.				
K511	<b>Utilities – Gas and Electric</b> Equipment using gas or related gas piping complies with NFPA 54, <i>National Fuel Gas Code</i> , electrical wiring and equipment complies with NFPA 70, <i>National Electric Code</i> . Existing installations can continue in service provided no hazard to life. 18.5.1.1, 19.5.1.1, 9.1.1, 9.1.2				
K521	<b>HVAC</b> Heating, ventilation, and air conditioning shall comply with 9.2 and shall be installed in accordance with the manufacturer's specifications. 18.5.2.1, 19.5.2.1, 9.2				
K522	<b>HVAC – Any Heating Device</b> Any heating device, other than a central heating plant, is designed and installed so combustible materials cannot be ignited by device, and has a safety feature to stop fuel and shut down equipment if there is excessive temperature or ignition failure. If fuel fired, the device also: <ul style="list-style-type: none"> <li>• is chimney or vent connected.</li> <li>• takes air for combustion from outside.</li> <li>• provides for a combustion system separate from occupied area atmosphere.</li> </ul> 18.5.2.2, 19.5.2.2				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K523	<b>HVAC – Suspended Unit Heaters</b> Suspended unit heaters are permitted provided the following are met: <ul style="list-style-type: none"> <li>• Not located in means of egress or in patient rooms.</li> <li>• Located high enough to be out of reach of people in the area.</li> <li>• Has a safety feature to stop fuel and shut down equipment if there is excessive temperature or ignition failure.</li> </ul> 18.5.2.3(1), 19.5.2.3(1)				
K524	<b>HVAC – Direct-Vent Gas Fireplaces</b> Direct-vent gas fireplaces, as defined in NFPA 54, inside of all smoke compartments containing patient sleeping areas comply with the requirements of 18.5.2.3(2), 19.5.2.3(2). 18.5.2.3(2), 19.5.2.3(2), NFPA 54				
K525	<b>HVAC – Solid Fuel-Burning Fireplaces</b> Solid fuel-burning fireplaces are permitted in other than patient sleeping areas provided: <ul style="list-style-type: none"> <li>• Areas are separated by 1-hour fire resistance construction.</li> <li>• Fireplace complies with 9.2.2.</li> <li>• Fireplace enclosure resists breakage up to 650°F and has heat-tempered glass.</li> <li>• Room has supervised CO detection per 9.8.</li> </ul> 18.5.2.3(3) and 19.5.2.3(3)				
K531	<b>Elevators</b> 2012 EXISTING Elevators comply with the provision of 9.4. Elevators are inspected and tested as specified in ASME A17.1, <i>Safety Code for Elevators and Escalators</i> . Firefighter's Service is operated monthly with a written record. Existing elevators conform to ASME/ANSI A17.3, <i>Safety Code for Existing Elevators and Escalators</i> . All existing elevators, having a travel distance of 25 feet or more above or below the level that best serves the needs of emergency personnel for firefighting purposes, conform with Firefighter's Service Requirements of ASME/ANSI A17.3. (Includes firefighter's service Phase I key recall and smoke detector automatic recall, firefighter's service Phase II emergency in-car key operation, machine room smoke detectors, and elevator lobby smoke detectors.) 19.5.3, 9.4.2, 9.4.3				



ID PREFIX		MET	NOT MET	N/A	REMARKS
K531	<p>2012 NEW</p> <p>Elevators comply with the provision of 9.4. Elevators are inspected and tested as specified in ASME A17.1, <i>Safety Code for Elevators and Escalators</i>. Firefighter's Service is operated monthly with a written record. New elevators conform to ASME/ANSI A17.1, <i>Safety Code for Elevators and Escalators</i>, including Firefighter's Service Requirements. (Includes firefighter's Phase I key recall and smoke detector automatic recall, firefighter's service Phase II emergency in-car key operation, machine room smoke detectors, and elevator lobby smoke detectors.)</p> <p>18.5.3, 9.4.2, 9.4.3</p>				
K532	<p><b>Escalators, Dumbwaiters, and Moving Walks</b></p> <p>2012 EXISTING</p> <p>Escalators, dumbwaiters, and moving walks comply with the provisions of 9.4.</p> <p>All existing escalators, dumbwaiters, and moving walks conform to the requirements of ASME/ANSI A17.3, <i>Safety Code for Existing Elevators and Escalators</i>.</p> <p>(Includes escalator emergency stop buttons and automatic skirt obstruction stop. For power dumbwaiters, includes hoistway door locking to keep doors closed except for floor where car is being loaded or unloaded.)</p> <p>19.5.3, 9.4.2.2</p>				
	<p>2012 NEW</p> <p>Escalators, dumbwaiters, and moving walks comply with the provisions of 9.4.</p> <p>18.5.3, 9.4.2.2</p>				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K541	<b>Rubbish Chutes, Incinerators, and Laundry Chutes</b> 2012 EXISTING (1) Any existing linen and trash chute, including pneumatic rubbish and linen systems, that opens directly onto any corridor shall be sealed by fire resistive construction to prevent further use or shall be provided with a fire door assembly having a fire protection rating of 1-hour. All new chutes shall comply with 9.5. (2) Any rubbish chute or linen chute, including pneumatic rubbish and linen systems, shall be provided with automatic extinguishing protection in accordance with 9.7. (3) Any trash chute shall discharge into a trash collection room used for no other purpose and protected in accordance with 8.4. (Existing laundry chutes permitted to discharge into same room are protected by automatic sprinklers in accordance with 19.3.5.9 or 19.3.5.7.) (4) Existing fuel-fed incinerators shall be sealed by fire resistive construction to prevent further use. 19.5.4, 9.5, 8.4, NFPA 82				
	2012 NEW Rubbish chutes, incinerators, and laundry chutes shall comply with the provisions of Section 9.5, unless otherwise specified in 18.5.4.2. <ul style="list-style-type: none"> <li>The fire resistance rating of chute charging room shall not be required to exceed 1-hour.</li> <li>Any rubbish chute or linen chute shall be provided with automatic extinguishing protection in accordance with Section 9.7.</li> <li>Chutes shall discharge into a trash collection room used for no other purpose and shall be protected in accordance with 8.7.</li> </ul> 18.5.4.2, 8.7, 9.5, 9.7, NFPA 82				
	<b>SECTION 6 – RESERVED</b>				
	<b>SECTION 7 – OPERATING FEATURES</b>				
K700	<b>Operating Features – Other</b> List in the REMARKS section any LSC Section 18.7 and 19.7 Operating Features requirements that are not addressed by the provided K-tags, but are deficient. This information, along with the applicable Life Safety Code or NFPA standard citation, should be included in Form CMS-2567.				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K711	<b>Evacuation and Relocation Plan</b> There is a written plan for the protection of all patients and for their evacuation in the event of an emergency. Employees are periodically instructed and kept informed with their duties under the plan, and a copy of the plan is readily available with telephone operator or with security. The plan addresses the basic response required of staff per 18/19.7.2.1.2 and provides for all of the fire safety plan components per 18/19.7.2.2. 18.7.1.1 through 18.7.1.3, 18.7.2.1.2, 18.7.2.2, 18.7.2.3, 19.7.1.1 through 19.7.1.3, 19.7.2.1.2, 19.7.2.2, 19.7.2.3				
K712	<b>Fire Drills</b> Fire drills include the transmission of a fire alarm signal and simulation of emergency fire conditions. Fire drills are held at expected and unexpected times under varying conditions, at least quarterly on each shift. The staff is familiar with procedures and is aware that drills are part of established routine. Where drills are conducted between 9:00 PM and 6:00 AM, a coded announcement may be used instead of audible alarms. 18.7.1.4 through 18.7.1.7, 19.7.1.4 through 19.7.1.7				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K741	<b>Smoking Regulations</b> Smoking regulations shall be adopted and shall include not less than the following provisions: (1) Smoking shall be prohibited in any room, ward, or compartment where flammable liquids, combustible gases, or oxygen is used or stored and in any other hazardous location, and such area shall be posted with signs that read NO SMOKING or shall be posted with the international symbol for no smoking. (2) In health care occupancies where smoking is prohibited and signs are prominently placed at all major entrances, secondary signs with language that prohibits smoking shall not be required. (3) Smoking by patients classified as not responsible shall be prohibited. (4) The requirement of 18.7.4(3) shall not apply where the patient is under direct supervision. (5) Ashtrays of noncombustible material and safe design shall be provided in all areas where smoking is permitted. (6) Metal containers with self-closing cover devices into which ashtrays can be emptied shall be readily available to all areas where smoking is permitted. 18.7.4, 19.7.4				
K751	<b>Draperies, Curtains, and Loosely Hanging Fabrics</b> Draperies, curtains including cubicle curtains and loosely hanging fabric or films shall be in accordance with 10.3.1. Excluding curtains and draperies: at showers and baths; on windows in patient sleeping room located in sprinklered compartments; and in non-patient sleeping rooms in sprinklered compartments where individual drapery or curtain panels do not exceed 48 square feet or total area does not exceed 20 percent of the wall. 18.7.5.1, 18.3.5.11, 19.7.5.1, 19.3.5.11, 10.3.1				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K752	<b>Upholstered Furniture and Mattresses</b> Newly introduced upholstered furniture meets Class I or char length, and heat release criteria in accordance with 10.3.2.1 and 10.3.3, unless the building is fully sprinklered. Newly introduced mattresses shall meet char length and heat release criteria in accordance with 10.3.2.2 and 10.3.4, unless the building is fully sprinklered. Upholstered furniture and mattresses belonging to nursing home residents do not have to meet these requirements as all nursing homes are required to be fully sprinklered. Newly introduced upholstered furniture and mattresses means purchased on or after the LSC final rule effective date. 18.7.5.2, 18.7.5.4, 19.7.5.2, 19.7.5.4				
K753	<b>Combustible Decorations</b> Combustible decorations shall be prohibited unless one of the following is met: <ul style="list-style-type: none"> <li>• Flame retardant or treated with approved fire-retardant coating that is listed and labeled for product.</li> <li>• Decorations meet NFPA 701.</li> <li>• Decorations exhibit heat release less than 100 kilowatts in accordance with NFPA 289.</li> <li>• Decorations, such as photographs, paintings and other art are attached to the walls, ceilings and non-fire-rated doors in accordance with 18.7.5.6(4) or 19.7.5.6(4).</li> <li>• The decorations in existing occupancies are in such limited quantities that a hazard of fire development or spread is not present.</li> </ul> 18.7.5.6, 19.7.5.6				
K761	<b>Maintenance, Inspection &amp; Testing - Doors</b> Fire doors assemblies are inspected and tested annually in accordance with NFPA 80 <i>Standard for Fire Doors and Other Opening Protectives</i> . Fire doors that are not located in required fire barriers, including corridor doors to patient rooms and smoke barrier doors, are routinely inspected as part of the facility maintenance program. Individuals performing the door inspection and testing have an understanding of the operating components of the doors. Written records of inspection and testing are maintained and are available for review. 18.7.6, 19.7.6, 8.3.3.1 (LSC), 5.2, 5.2.3 (NFPA 80)				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K754	<b>Soiled Linen and Trash Containers</b> Soiled linen or trash collection receptacles shall not exceed 32 gallons in capacity. The average density of container capacity in a room or space shall not exceed 0.5 gallons/square feet. A total container capacity of 32 gallons shall not be exceeded within any 64 square feet area. Mobile soiled linen or trash collection receptacles with capacities greater than 32 gallons shall be located in a room protected as a hazardous area when not attended. Containers used solely for recycling are permitted to be excluded from the above requirements where each container is $\leq$ 96 gal. unless attended, and containers for combustibles are labeled and listed as meeting FM Approval Standard 6921 or equivalent. 18.7.5.7, 19.7.5.7				
K771	<b>Engineer Smoke Control Systems</b> 2012 EXISTING When installed, engineered smoke control systems are tested in accordance with established engineering principles. Test documentation is maintained on the premises. 19.7.7				
	2012 NEW When installed, engineered smoke control systems are tested in accordance with NFPA 92, <i>Standard for Smoke Control Systems</i> . Test documentation is maintained on the premises. 18.7.7				
K781	<b>Portable Space Heaters</b> Portable space heating devices shall be prohibited in all health care occupancies. Unless used in nonsleeping staff and employee areas where the heating elements do not exceed 212 degrees Fahrenheit (100 degrees Celsius). 18.7.8, 19.7.8				
K791	<b>Construction, Repair, and Improvement Operations</b> Construction, repair, and improvement operations shall comply with 4.6.10. Any means of egress in any area undergoing construction, repair, or improvements shall be inspected daily to ensure its ability to be used instantly in case of emergency and compliance with NFPA 241. 18.7.9, 19.7.9, 4.6.10, 7.1.10.1				

ID PREFIX		MET	NOT MET	N/A	REMARKS
	<b>PART II – HEALTH CARE FACILITIES CODE REQUIREMENTS</b>				
K900	<b>Health Care Facilities Code - Other</b> List in the REMARKS section any NFPA 99 requirements (excluding Chapter 7, 8, 12, and 13) that are not addressed by the provided K-Tags, but are deficient. This information, along with the applicable Health Care Facilities Code or NFPA standard citation, should be included on Form CMS-2567.				
K901	<b>Fundamentals – Building System Categories</b> Building systems are designed to meet Category 1 through 4 requirements as detailed in NFPA 99. Categories are determined by a formal and documented risk assessment procedure performed by qualified personnel. Chapter 4 (NFPA 99)				
K902	<b>Gas and Vacuum Piped Systems – Other</b> List in the REMARKS section any NFPA 99 Chapter 5 Gas and Vacuum Systems requirements that are not addressed by the provided K-Tags, but are deficient. This information, along with the applicable Life Safety Code or NFPA standard citation, should be included on Form CMS-2567. Chapter 5 (NFPA 99)				
K903	<b>Gas and Vacuum Piped Systems – Categories</b> Medical gas, medical air, surgical vacuum, WAGD, and air supply systems are designated: <input type="checkbox"/> Category 1. Systems in which failure is likely to cause major injury or death. <input type="checkbox"/> Category 2. Systems in which failure is likely to cause minor injury. <input type="checkbox"/> Category 3. Systems in which failure is not likely to cause injury, but can cause discomfort. Deep sedation and general anesthesia are not to be administered using a Category 3 medical gas system. 5.1.1.1, 5.2.1, 5.3.1.1, 5.3.1.5 (NFPA 99)				
K904	<b>Gas and Vacuum Piped Systems – Warning Systems</b> All master, area, and local alarm systems used for medical gas and vacuum systems comply with appropriate Category warning system requirements, as applicable. 5.1.9, 5.2.9, 5.3.6.2.2 (NFPA 99)				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K905	<b>Gas and Vacuum Piped Systems – Central Supply System Identification and Labeling</b> Containers, cylinders and tanks are designed, fabricated, tested, and marked in accordance with 5.1.3.1.1 through 5.1.3.1.7. Locations containing only oxygen or medical air have doors labeled with "Medical Gases, NO Smoking or Open Flame". Locations containing other gases have doors labeled "Positive Pressure Gases, NO Smoking or Open Flame, Room May Have Insufficient Oxygen, Open Door and Allow Room to Ventilate Before Opening." 5.1.3.1, 5.2.3.1, 5.3.10 (NFPA 99)				
K906	<b>Gas and Vacuum Piped Systems – Central Supply System Operations</b> Adaptors or conversion fittings are prohibited. Cylinders are handled in accordance with 11.6.2. Only cylinders, reusable shipping containers, and their accessories are stored in rooms containing central supply systems or cylinders. No flammable materials are stored with cylinders. Cryogenic liquid storage units intended to supply the facility are not used to transfill. Cylinders are kept away from sources of heat. Valve protection caps are secured in place, if supplied, unless cylinder is in use. Cylinders are not stored in tightly closed spaces. Cylinders in use and storage are prevented from exceeding 130°F, and nitrous oxide and carbon dioxide cylinders are prevented from reaching temperatures lower than manufacture recommendations or 20°F. Full or empty cylinders, when not connected, are stored in locations complying with 5.1.3.3.2 through 5.1.3.3.3, and are not stored in enclosures containing motor-driven machinery, unless for instrument air reserve headers. 5.1.3.2, 5.1.3.3.17, 5.1.3.3.1.8, 5.1.3.3.4, 5.2.3.2, 5.2.3.3, 5.3.6.20.4, 5.6.20.5, 5.3.6.20.7, 5.3.6.20.8, 5.3.6.20.9 (NFPA 99)				
K907	<b>Gas and Vacuum Piped Systems – Maintenance Program</b> Medical gas, vacuum, WAGD, or support gas systems have documented maintenance programs. The program includes an inventory of all source systems, control valves, alarms, manufactured assemblies, and outlets. Inspection and maintenance schedules are established through risk assessment considering manufacturer recommendations. Inspection procedures and testing methods are established through risk assessment. Persons maintaining systems are qualified as demonstrated by training and certification or credentialing to the requirements of AASE 6030 or 6040. 5.1.14.2.1, 5.1.14.2.2, 5.1.15, 5.2.14, 5.3.13.4.2 (NFPA 99)				



ID PREFIX		MET	NOT MET	N/A	REMARKS
K908	<b>Gas and Vacuum Piped Systems – Inspection and Testing Operations</b> The gas and vacuum systems are inspected and tested as part of a maintenance program and include the required elements. Records of the inspections and testing are maintained as required. 5.1.14.2.3, B.5.2, 5.2.13, 5.3.13, 5.3.13.4 (NFPA 99)				
K909	<b>Gas and Vacuum Piped Systems – Information and Warning Signs</b> Piping is labeled by stencil or adhesive markers identifying the gas or vacuum system, including the name of system or chemical symbol, color code (Table 5.1.11), and operating pressure if other than standard. Labels are at intervals not more than 20 feet, are in every room, at both sides of wall penetrations, and on every story traversed by riser. Piping is not painted. Shutoff valves are identified with the name or chemical symbol of the gas or vacuum system, room or area served, and caution to not use the valve except in emergency. 5.1.14.3, 5.1.11.1, 5.1.11.2, 5.2.11, 5.3.13.3, 5.3.11 (NFPA 99)				
K910	<b>Gas and Vacuum Piped Systems – Modifications</b> Whenever modifications are made that breach the pipeline, any necessary installer and verification test specified in 5.1.2 is conducted on the downstream portion of the medical gas piping system. Permanent records of all tests required by system verification tests are maintained. 5.1.14.4.1, 5.1.14.4.6, 5.2.13, 5.3.13.4.3 (NFPA 99)				
K911	<b>Electrical Systems – Other</b> List in the REMARKS section any NFPA 99 Chapter 6 Electrical Systems requirements that are not addressed by the provided K-Tags, but are deficient. This information, along with the applicable Life Safety Code or NFPA standard citation, should be included on Form CMS-2567. Chapter 6 (NFPA 99)				
K912	<b>Electrical Systems – Receptacles</b> Power receptacles have at least one, separate, highly dependable grounding pole capable of maintaining low-contact resistance with its mating plug. In pediatric locations, receptacles in patient rooms, bathrooms, play rooms, and activity rooms, other than nurseries, are listed tamper-resistant or employ a listed cover. If used in patient care room, ground-fault circuit interrupters (GFCI) are listed. 6.3.2.2.6.2 (F), 6.3.2.2.4.2 (NFPA 99)				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K913	<b>Electrical Systems – Wet Procedure Locations</b> Operating rooms are considered wet procedure locations, unless otherwise determined by a risk assessment conducted by the facility governing body. Operating rooms defined as wet locations are protected by either isolated power or ground-fault circuit interrupters. A written record of the risk assessment is maintained and available for inspection. 6.3.2.2.8.4, 6.3.2.2.8.7, 6.4.4.2				
K914	<b>Electrical Systems – Maintenance and Testing</b> Hospital-grade receptacles at patient bed locations and where deep sedation or general anesthesia is administered, are tested after initial installation, replacement or servicing. Additional testing is performed at intervals defined by documented performance data. Receptacles not listed as hospital-grade at these locations are tested at intervals not exceeding 12 months. Line isolation monitors (LIM), if installed, are tested at intervals of ≤ 1 month by actuating the LIM test switch per 6.3.2.6.3.6, which activates both visual and audible alarm. For LIM circuits with automated self-testing, this manual test is performed at intervals ≤ 12 months. LIM circuits are tested per 6.3.3.3.2 after any repair or renovation to the electric distribution system. Records are maintained of required tests and associated repairs or modifications, containing date, room or area tested, and results. 6.3.4 (NFPA 99)				
K915	<b>Electrical Systems – Essential Electric System Categories</b> <input type="checkbox"/> Critical care rooms (Category 1) in which electrical system failure is likely to cause major injury or death of patients, including all rooms where electric life support equipment is required, are served by a Type 1 EES. <input type="checkbox"/> General care rooms (Category 2) in which electrical system failure is likely to cause minor injury to patients (Category 2) are served by a Type 1 or Type 2 EES. <input type="checkbox"/> Basic care rooms (Category 3) in which electrical system failure is not likely to cause injury to patients and rooms other than patient care rooms are not required to be served by an EES. Type 3 EES life safety branch has an alternate source of power that will be effective for 1 1/2 hours. 3.3.138, 6.3.2.2.10, 6.6.2.2.2, 6.6.3.1.1 (NFPA 99), TIA 12-3				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K916	<b>Electrical Systems – Essential Electric System Alarm Annunciator</b> A remote annunciator that is storage battery powered is provided to operate outside of the generating room in a location readily observed by operating personnel. The annunciator is hard-wired to indicate alarm conditions of the emergency power source. A centralized computer system (e.g., building information system) is not to be substituted for the alarm annunciator. 6.4.1.1.17, 6.4.1.1.17.5 (NFPA 99)				
K917	<b>Electrical Systems – Essential Electric System Receptacles</b> Electrical receptacles or cover plates supplied from the life safety and critical branches have a distinctive color or marking. 6.4.2.2.6, 6.5.2.2.4.2, 6.6.2.2.3.2 (NFPA 99)				
K918	<b>Electrical Systems – Essential Electric System Maintenance and Testing</b> The generator or other alternate power source and associated equipment is capable of supplying service within 10 seconds. If the 10-second criterion is not met during the monthly test, a process shall be provided to annually confirm this capability for the life safety and critical branches. Maintenance and testing of the generator and transfer switches are performed in accordance with NFPA 110. Generator sets are inspected weekly, exercised under load 30 minutes 12 times a year in 20-40 day intervals, and exercised once every 36 months for 4 continuous hours. Scheduled test under load conditions include a complete simulated cold start and automatic or manual transfer of all EES loads, and are conducted by competent personnel. Maintenance and testing of stored energy power sources (Type 3 EES) are in accordance with NFPA 111. Main and feeder circuit breakers are inspected annually, and a program for periodically exercising the components is established according to manufacturer requirements. Written records of maintenance and testing are maintained and readily available. EES electrical panels and circuits are marked, readily identifiable, and separate from normal power circuits. Minimizing the possibility of damage of the emergency power source is a design consideration for new installations. 6.4.4, 6.5.4, 6.6.4 (NFPA 99), NFPA 110, NFPA 111, 700.10 (NFPA 70)				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K919	<b>Electrical Equipment – Other</b> List in the REMARKS section any NFPA 99 Chapter 10, <i>Electrical Equipment</i> , requirements that are not addressed by the provided K-Tags, but are deficient. This information, along with the applicable Life Safety Code or NFPA standard citation, should be included on Form CMS-2567. Chapter 10 (NFPA 99)				
K920	<b>Electrical Equipment – Power Cords and Extension Cords</b> Power strips in a patient care vicinity are only used for components of movable patient-care-related electrical equipment (PCREE) assemblies that have been assembled by qualified personnel and meet the conditions of 10.2.3.6. Power strips in the patient care vicinity may not be used for non-PCREE (e.g., personal electronics), except in long-term care resident rooms that do not use PCREE. Power strips for PCREE meet UL 1363A or UL 60601-1. Power strips for non-PCREE in the patient care rooms (outside of vicinity) meet UL 1363. In non-patient care rooms, power strips meet other UL standards. All power strips are used with general precautions. Extension cords are not used as a substitute for fixed wiring of a structure. Extension cords used temporarily are removed immediately upon completion of the purpose for which it was installed and meets the conditions of 10.2.4. 10.2.3.6 (NFPA 99), 10.2.4 (NFPA 99), 400-8 (NFPA 70), 590.3(D) (NFPA 70), TIA 12-5				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K921	<b>Electrical Equipment – Testing and Maintenance Requirements</b> The physical integrity, resistance, leakage current, and touch current tests for fixed and portable patient-care related electrical equipment (PCREE) is performed as required in 10.3. Testing intervals are established with policies and protocols. All PCREE used in patient care rooms is tested in accordance with 10.3.5.4 or 10.3.6 before being put into service and after any repair or modification. Any system consisting of several electrical appliances demonstrates compliance with NFPA 99 as a complete system. Service manuals, instructions, and procedures provided by the manufacturer include information as required by 10.5.3.1.1 and are considered in the development of a program for electrical equipment maintenance. Electrical equipment instructions and maintenance manuals are readily available, and safety labels and condensed operating instructions on the appliance are legible. A record of electrical equipment tests, repairs, and modifications is maintained for a period of time to demonstrate compliance in accordance with the facility's policy. Personnel responsible for the testing, maintenance and use of electrical appliances receive continuing training. 10.3, 10.5.2.1, 10.5.2.1.2, 10.5.2.5, 10.5.3, 10.5.6, 10.5.8				
K922	<b>Gas Equipment – Other</b> List in the REMARKS section any NFPA 99 Chapter 11 Gas Equipment requirements that are not addressed by the provided K-Tags, but are deficient. This information, along with the applicable Life Safety Code or NFPA standard citation, should be included on Form CMS-2567. Chapter 11 (NFPA 99)				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K923	<p><b>Gas Equipment – Cylinder and Container Storage</b></p> <p><b>≥ 3,000 cubic feet</b> Storage locations are designed, constructed, and ventilated in accordance with 5.1.3.3.2 and 5.1.3.3.3.</p> <p><b>&gt; 300 but &lt;3,000 cubic feet</b> Storage locations are outdoors in an enclosure or within an enclosed interior space of non- or limited- combustible construction, with door (or gates outdoors) that can be secured. Oxidizing gases are not stored with flammables, and are separated from combustibles by 20 feet (5 feet if sprinklered) or enclosed in a cabinet of noncombustible construction having a minimum 1/2 hr. fire protection rating.</p> <p><b>≤ 300 cubic feet</b> In a single smoke compartment, individual cylinders available for immediate use in patient care areas with an aggregate volume of ≤ 300 cubic feet are not required to be stored in an enclosure. Cylinders must be handled with precautions as specified in 11.6.2.</p> <p>A precautionary sign readable from 5 feet is on each door or gate of a cylinder storage room, where the sign includes the wording as a minimum "CAUTION: OXIDIZING GAS(ES) STORED WITHIN NO SMOKING".</p> <p>Storage is planned so cylinders are used in order of which they are received from the supplier. Empty cylinders are segregated from full cylinders. When facility employs cylinders with integral pressure gauge, a threshold pressure considered empty is established. Empty cylinders are marked to avoid confusion. Cylinders stored in the open are protected from weather.</p> <p>11.3.1, 11.3.2, 11.3.3, 11.3.4, 11.6.5 (NFPA 99)</p>				
K924	<p><b>Gas Equipment – Testing and Maintenance Requirements</b></p> <p>Anesthesia apparatus are tested at the final path to patient after any adjustment, modification or repair. Before the apparatus is returned to service, each connection is checked to verify proper gas and an oxygen analyzer is used to verify oxygen concentration. Defective equipment is immediately removed from service. Areas designated for servicing of oxygen equipment are clean and free of oil, grease, or other flammables. Manufacturer service manuals are used to maintain equipment and a scheduled maintenance program is followed.</p> <p>11.4.1.3, 11.5.1.3, 11.6.2.5, 11.6.2.6 (NFPA 99)</p>				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K925	<b>Gas Equipment – Respiratory Therapy Sources of Ignition</b> Smoking materials are removed from patients receiving respiratory therapy. When a nasal cannula is delivering oxygen outside of a patient's room, no sources of ignition are within in the site of intentional expulsion (1-foot). When other oxygen deliver equipment is used or oxygen is delivered inside a patient's room, no sources of ignition are within the area are of administration (15-feet). Solid fuel-burning appliances is not in the area of administration. Nonmedical appliances with hot surfaces or sparking mechanisms are not within oxygen-delivery equipment or site of intentional expulsion. 11.5.1.1, TIA 12-6 (NFPA 99)				
K926	<b>Gas Equipment – Qualifications and Training of Personnel</b> Personnel concerned with the application, maintenance and handling of medical gases and cylinders are trained on the risk. Facilities provide continuing education, including safety guidelines and usage requirements. Equipment is serviced only by personnel trained in the maintenance and operation of equipment. 11.5.2.1 (NFPA 99)				
K927	<b>Gas Equipment – Transfilling Cylinders</b> Transfilling of oxygen from one cylinder to another is in accordance with CGA P-2.5, <i>Transfilling of High Pressure Gaseous Oxygen Used for Respiration</i> . Transfilling of any gas from one cylinder to another is prohibited in patient care rooms. Transfilling to liquid oxygen containers or to portable containers over 50 psi comply with conditions under 11.5.2.3.1 (NFPA 99). Transfilling to liquid oxygen containers or to portable containers under 50 psi comply with conditions under 11.5.2.3.2 (NFPA 99). 11.5.2.2 (NFPA 99)				

ID PREFIX		MET	NOT MET	N/A	REMARKS
K928	<b>Gas Equipment – Labeling Equipment and Cylinders</b> Equipment listed for use in oxygen-enriched atmospheres are so labeled. Oxygen metering equipment and pressure reducing regulators are labeled "OXYGEN-USE NO OIL". Flowmeters, pressure reducing regulators, and oxygen-dispensing apparatus are clearly and permanently labeled designating the gases for which they are intended. Oxygen-metering equipment, pressure reducing regulators, humidifiers, and nebulizers are labeled with name of manufacturer or supplier. Cylinders and containers are labeled in accordance with CGA C-7. Color coding is not utilized as the primary method of determining cylinder or container contents. All labeling is durable and withstands cleaning or disinfecting. 11.5.3.1 (NFPA 99)				
K929	<b>Gas Equipment – Precautions for Handling Oxygen Cylinders and Manifolds</b> Handling of oxygen cylinders and manifolds is based on CGA G-4, Oxygen. Oxygen cylinders, containers, and associated equipment are protected from contact with oil and grease, from contamination, protected from damage, and handled with care in accordance with precautions provided under 11.6.2.1 through 11.6.2.4 (NFPA 99). 11.6.2 (NFPA 99)				
K930	<b>Gas Equipment – Liquid Oxygen Equipment</b> The storage and use of liquid oxygen in base reservoir containers and portable containers comply with sections 11.7.2 through 11.7.4 (NFPA 99). 11.7 (NFPA 99)				
K931	<b>Hyperbaric Facilities</b> All occupancies containing hyperbaric facilities comply with construction, equipment, administration, and maintenance requirements of NFPA 99. Chapter 14 (NFPA 99)				
K932	<b>Features of Fire Protection – Other</b> List in the REMARKS section any NFPA 99 Chapter 15 Features of Fire Protection requirements that are not addressed by the provided K-Tags, but are deficient. This information, along with the applicable Life Safety Code or NFPA standard citation, should be included on Form CMS-2567. Chapter 15 (NFPA 99)				



ID PREFIX		MET	NOT MET	N/A	REMARKS
K933	<p><b>Features of Fire Protection – Fire Loss Prevention in Operating Rooms</b></p> <p>Periodic evaluations are made of hazards that could be encountered during surgical procedures, and fire prevention procedures are established. When flammable germicides or antiseptics are employed during surgeries utilizing electrosurgery, cautery or lasers:</p> <ul style="list-style-type: none"> <li>• packaging is non-flammable.</li> <li>• applicators are in unit doses.</li> <li>• Preoperative "time-out" is conducted prior the initiation of any surgical procedure to verify: <ul style="list-style-type: none"> <li>○ application site is dry prior to draping and use of surgical equipment.</li> <li>○ pooling of solution has not occurred or has been corrected.</li> <li>○ solution-soaked materials have been removed from the OR prior to draping and use of surgical devices.</li> <li>○ policies and procedures are established outlining safety precautions related to the use of flammable germicide or antiseptic use.</li> </ul> </li> </ul> <p>Procedures are established for operating room emergencies including alarm activation, evacuation, equipment shutdown, and control operations. Emergency procedures include the control of chemical spills, and extinguishment of drapery, clothing and equipment fires. Training is provided to new OR personnel (including surgeons), continuing education is provided, incidents are reviewed monthly, and procedures are reviewed annually.</p> <p>15.13 (NFPA 99)</p>				

**PART III – RECOMMENDATION FOR WAIVER OF SPECIFIC LIFE SAFETY CODE PROVISIONS**

For each item of the Life Safety Code recommended for waiver, list the survey report form item number and state the reason for the conclusion that: (a) the specific provisions of the code, if rigidly applied, would result in unreasonable hardship on the facility, and (b) the waiver of such unmet provisions will not adversely affect the health and safety of the patients. If additional space is required, attach additional sheet(s).

**PROVISION NUMBER(S)****JUSTIFICATION**

K400

Surveyor ( <i>Signature</i> )	Title	Office	Date
Fire Authority Official ( <i>Signature</i> )	Title	Office	Date

Provider Number	Facility Name	Survey Date
K1		*K4

K6	DATE OF PLAN APPROVAL	K3	MULTIPLE CONSTRUCTION	<input type="checkbox"/>	A. BUILDING
			TOTAL NUMBER OF BUILDINGS _____		B. WING
			NUMBER OF THIS BUILDING _____		C. FLOOR
					D. APARTMENT UNIT

HEALTH CARE FORM		
12	2786R	2012 EXISTING
13	2786R	2012 NEW

AHCO FORM		
14	2786U	2012 EXISTING
15	2786U	2012 NEW

ICF/IID FORM		
16	2786V, W, X	2012 EXISTING
17	2786V, W, X	2012 NEW

\*K7

SELECT NUMBER OF FORM USED FROM ABOVE

*(Check if K321 or K351 are marked as not applicable in the 2786 M, R, T, U, V, W, X, and Y.)*

K321:

K351:

\*K9 FACILITY MEETS LSC BASED ON *(Check all that Apply)*

A1. <input type="checkbox"/>	A2. <input type="checkbox"/>	A3. <input type="checkbox"/>	A4. <input type="checkbox"/>	A5. <input type="checkbox"/>
(COMP. WITH ALL PROVISIONS)	(ACCEPTABLE POC)	(WAIVERS)	(FSSES)	(PERFORMANCE BASED DESIGN)

<p>FACILITY DOES NOT MEET LSC</p>	<p>K0180</p>		
	<p>B. <input type="checkbox"/></p>	<p>A. <input type="checkbox"/></p> <p><b>FULLY SPRINKLERED</b> (All required areas are sprinklered)</p>	<p>B. <input type="checkbox"/></p> <p><b>PARTIALLY SPRINKLERED</b> (Not all required areas are sprinklered)</p>

Form CMS-2786R (07/2018)

**FIRE SAFETY SURVEY REPORT  
CRUCIAL DATA EXTRACT  
(TO BE USED WITH CMS-2786 FORMS)**

PROVIDER NUMBER  K1 245496	FACILITY NAME  <b>MINNEOTA MANOR HEALTH CARE CENTER</b>	SURVEY DATE  <b>*K4 09/16/2021</b>
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K6 DATE OF PLAN APPROVAL	<div style="display: flex; justify-content: space-between;"> <div> K3 : MULTIPLE CONSTRUCTION   TOTAL NUMBER OF BUILDINGS <u>1</u>   NUMBER OF THIS BUILDING <u>01</u> </div> <div style="border: 1px solid black; padding: 2px; text-align: center;">A</div> <div> A BUILDING  B WING  C FLOOR  D APARTMENT UNIT </div> </div>
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LSC FORM INDICATOR  <table border="1" style="width: 100%; border-collapse: collapse; margin-bottom: 5px;"> <tr><th colspan="3">Health Care Form</th></tr> <tr><td style="width: 10%;">12</td><td style="width: 60%;">2786 R</td><td style="width: 30%;">2012 EXISTING</td></tr> <tr><td>13</td><td>2786 R</td><td>2012 NEW</td></tr> </table> <table border="1" style="width: 100%; border-collapse: collapse; margin-bottom: 5px;"> <tr><th colspan="3">ASC Form</th></tr> <tr><td style="width: 10%;">14</td><td style="width: 60%;">2786 U</td><td style="width: 30%;">2012 EXISTING</td></tr> <tr><td>15</td><td>2786 U</td><td>2012 NEW</td></tr> </table> <table border="1" style="width: 100%; border-collapse: collapse;"> <tr><th colspan="3">ICF/MR Form</th></tr> <tr><td style="width: 10%;">16</td><td style="width: 60%;">2786 V, W, X</td><td style="width: 30%;">2012 EXISTING</td></tr> <tr><td>17</td><td>2786 V, W, X</td><td>2012 NEW</td></tr> </table> *K7 <span style="border: 1px solid black; padding: 0 5px;">12</span> SELECT NUMBER OF FORM USED FROM ABOVE  <i>(Check if K321 or K351 are marked as not applicable in the 2786 M, R, T, U, V, W, X, Y and Z.)</i>  <div style="display: flex; justify-content: space-around;"> <div>K321: <span style="border: 1px solid black; padding: 0 10px;">3</span></div> <div>K351: <span style="border: 1px solid black; padding: 0 10px;">3</span></div> </div>	Health Care Form			12	2786 R	2012 EXISTING	13	2786 R	2012 NEW	ASC Form			14	2786 U	2012 EXISTING	15	2786 U	2012 NEW	ICF/MR Form			16	2786 V, W, X	2012 EXISTING	17	2786 V, W, X	2012 NEW	COMPLETE IF ICF/MR IS SURVEYED UNDER CHAPTER 21 SMALL (16 BEDS OR LESS)  <div style="display: flex; justify-content: space-between;"> K8: <span style="border: 1px solid black; padding: 0 10px;"></span> <div> 1 PROMPT  2 SLOW  3 IMPRACTICAL </div> </div> <hr/> LARGE  <div style="display: flex; justify-content: space-between;"> K8: <span style="border: 1px solid black; padding: 0 10px;"></span> <div> 4 PROMPT  5 SLOW  6 IMPRACTICAL </div> </div> <hr/> APARTMENT HOUSE  <div style="display: flex; justify-content: space-between;"> K8: <span style="border: 1px solid black; padding: 0 10px;"></span> <div> 7 PROMPT  8 SLOW  9 IMPRACTICAL </div> </div> <hr/> ENTER E-SCORE HERE  <div style="display: flex; justify-content: space-between;"> K5: <span style="border: 1px solid black; padding: 0 10px;"></span> <div>e.g 2.5</div> </div>
Health Care Form																												
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17	2786 V, W, X	2012 NEW																										

\*K9 : FACILITY MEETS LSC BASED ON: *(Check all that apply)*

A1   
 (COMP. WITH ALL PROVISIONS)

A2 X  
 (ACCEPTABLE POC)

A3   
 (WAIVERS)

A4   
 (FSSES)

A5   
 (PERFORMANCE BASED DESIGN)

FACILITY DOES NOT MEET LSC:  B. <span style="border: 1px solid black; padding: 0 10px;"></span>	K180: <div style="display: flex; justify-content: space-around; align-items: flex-end; margin-top: 10px;"> <div style="text-align: center;"> A. <span style="border: 1px solid black; padding: 0 10px;">X</span>            FULLY SPRINKLERED            (All required areas are sprinklered)         </div> <div style="text-align: center;"> B. <span style="border: 1px solid black; padding: 0 10px;"></span>            PARTIALLY SPRINKLERED            (Not all required areas are sprinklered)         </div> <div style="text-align: center;"> C. <span style="border: 1px solid black; padding: 0 10px;"></span>            NONE            (No sprinkler system)         </div> </div>
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\*MANDATORY