

Department of Health & Human Services
Centers for Medicare & Medicaid Services

Printed: 07/30/2026
Form Approved OMB
No. 0938-0391

STATEMENT OF DEFICIENCIES AND PLAN OF CORRECTION	(X1) PROVIDER/SUPPLIER/CLIA IDENTIFICATION NUMBER: 085054	(X2) MULTIPLE CONSTRUCTION A. Building B. Wing	(X3) DATE SURVEY COMPLETED 05/15/2026
NAME OF PROVIDER OR SUPPLIER Cadia Rehabilitation Pike Creek		STREET ADDRESS, CITY, STATE, ZIP CODE 3540 Three Little Bakers Blvd Wilmington, DE 19808	
For information on the nursing home's plan to correct this deficiency, please contact the nursing home or the state survey agency.			
(X4) ID PREFIX TAG	SUMMARY STATEMENT OF DEFICIENCIES (Each deficiency must be preceded by full regulatory or LSC identifying information)		
F 0686 Level of Harm - Minimal harm or potential for actual harm Residents Affected - Few	Provide appropriate pressure ulcer care and prevent new ulcers from developing. **NOTE- TERMS IN BRACKETS HAVE BEEN EDITED TO PROTECT CONFIDENTIALITY** Based on interview, record review, and review of the facility's policy, the facility failed to ensure ordered pressure ulcer treatments were completed and/or newly identified skin changes were reported for three of five sampled residents (Resident (R) 182, R37, and R20) reviewed for pressure ulcers out of a total sample of 55. Failure to complete wound treatments as ordered and report changes in skin integrity placed the residents at increased risk of infection and wound deterioration. Findings include: Findings include: 1. Review of R182's admission Record, located under the Profile tab of the electronic medical record (EMR), revealed R182 was admitted to the facility on [DATE]. R182's pertinent diagnoses included spinal muscular atrophy, pneumonia, acute and chronic respiratory failure with hypoxia, severe sepsis with septic shock, severe protein-calorie malnutrition, hypotension, polyneuropathy, and generalized muscle weakness. Review of R182's 5-day Minimum Data Set (MDS), with an Assessment Reference Date (ARD) of 10/26/25 and located under the MDS tab of the EMR, revealed a Brief Interview for Mental Status (BIMS) score of 13 out of 15, which indicated the resident was cognitively intact. The resident did not exhibit behaviors or rejection of care during the review period. R182 required total dependence on staff for bed mobility, transfers, bathing, toileting, and toileting hygiene, and required substantial to maximum assistance with eating and oral hygiene. R182 was always incontinent with bowel and bladder. R182 was at risk for pressure ulcer development and had three unstageable pressure ulcers present on admission. R182 utilized pressure-reducing devices for both the bed and wheelchair and received pressure ulcer/injury care during the review period. Review of R182's Care Plan, dated 10/31/25 and located under the Care Plan tab of the EMR, revealed R182 had unstageable pressure ulcers to his left buttock, right buttock, and right great toe that were present upon admission and a reddened area to the sacrum. The pertinent interventions directed staff to administer treatments as ordered and monitor for effectiveness. Review of R182's Physician Orders, dated 10/2025 through 11/2025, located under the Orders tab of the EMR, revealed the following pertinent orders: -10/21/25: Santyl External Ointment 250 Unit/ gram (GM) (collagenase). Apply to bilateral buttocks topical one time a day for wound maintenance. Discontinued 11/12/25. -10/22/25: Cleanse left buttock with normal saline, pat dry, apply Santyl, cover with calcium alginate, and dry dressing one time day for wound maintenance. Discontinued 11/12/25. -10/22/25: Cleanse right buttock with normal saline, pat dry, apply Santyl and cover with calcium (continued on next page)		

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.

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

TITLE

(X6) DATE

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F 0686 Level of Harm - Minimal harm or potential for actual harm Residents Affected - Few	Review of R20's quarterly MDS, with an ARD of 05/01/26, and located in Internet Quality Improvement and Evaluation System (IQIES), revealed R20 had a BIMS score of 00 out of 15, which indicated the resident was severely cognitively impaired. It was recorded R20 was at high risk for the development of pressure ulcers and had no current skin concerns. It was recorded R20 was dependent on staff for all activities of daily living (ADLs) and required a two-person assist with a Hoyer lift for transfers. Review of the R20's Care Plan, revised on 01/28/26, and located under the Care Plan tab in the EMR, revealed interventions to prevent skin breakdown and promote skin integrity. Interventions included completion of Braden risk assessments, a wheelchair cushion, pressure redistribution mattress, offloading heels while in bed, and skin checks every two hours and as needed. During an interview on 05/14/26 at 1:35 PM, R20's family member (F) 1 stated after assisting with incontinence care earlier in the day (on 05/14/26), she observed a discoloration of the skin on R20's right upper buttocks. During an interview on 05/14/26 at 1:57 PM, Certified Nursing Assistant (CNA) 13 stated he was assigned to R20 and provided incontinence care at approximately 8:30 AM and again at 1:02 PM. CNA13 revealed he observed a red area on the resident's right buttock at 8:30 AM. CNA13 stated that the area appeared unchanged in size and condition during the subsequent care provided at 1:02 PM. CNA13 stated he did not report the identified new red area on the R20's right buttock to his charge nurse, as directed by facility protocol. During an interview on 05/14/26 at 4:24 PM, Unit Manager (UM) 4 confirmed CNAs were expected to report any observed skin changes. UM4 stated R20's newly identified skin change was identified at approximately 9:00 AM on 05/14/26. UM4 reviewed R20's Kardex and stated the documentation indicated no skin concerns were observed or noted. Review of a Progress Note, dated 05/15/26 at 11:57 AM and located under the Progress Notes tab in the EMR, revealed R20 was noted to have an open area to the upper right buttock measuring approximately 0.5 centimeters (cm). During an interview on 05/15/26 at 5:45 PM, the Administrator confirmed staff should adhere to ordered care interventions to ensure early identification of changes in a resident's condition. Review of the facility's policy titled, Pressure Ulcer Prevention and Management, last revised 01/17/23, revealed, It is the policy of Cadia Healthcare to promote skin integrity in residents through the recognition, treatment and prevention of pressure ulcers . Residents will have a plan of care developed and updated as needed. Interventions implemented are added to the resident's plan of care .		

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F 0755 Level of Harm - Minimal harm or potential for actual harm Residents Affected - Few	Provide pharmaceutical services to meet the needs of each resident and employ or obtain the services of a licensed pharmacist. **NOTE- TERMS IN BRACKETS HAVE BEEN EDITED TO PROTECT CONFIDENTIALITY** Based on interview, record review, and facility policy review, the facility failed to safeguard controlled medications used in facility stock and failed to safeguard controlled medications and/or ensure the availability of physician ordered medications for two of 55 sampled residents (Resident (R)178 and R5). These failed practices placed residents at risk for medication diversion, unmanaged symptoms, and failure to receive medications as prescribed. Findings include:1. Review of a Pharmacy Delivery Manifest, dated 09/10/25, provided by the Director of Nursing (DON), revealed one pill of Oxycodone IR (an immediate release narcotic pain medication) 5 milligrams (mg) was delivered to the facility for a stock replacement order in the Omnicell (a secured automated medication dispensing cabinet). The Pharmacy Delivery Manifest for 09/10/25 had a delivery time of 7:28 AM and was signed by Licensed Practical Nurse (LPN)10. During an interview on 05/13/26 at 10:00 AM, the DON stated she received an email from the pharmacy on 09/15/25 informing the facility the Oxycodone IR 5 mg replacement pill for the Omnicell, delivered on 09/10/25, was never placed into the Omnicell. The DON stated she then checked the Omnicell, and the Oxycodone IR 5 mg pill was not in the Omnicell. The DON stated she then interviewed LPN10, who signed for the Oxycodone delivery on 9/10/25. The DON stated LPN10 verified she did accept and sign for the Oxycodone IR 5 mg pill from the pharmacy on 09/10/25. The DON stated she was told LPN10 and LPN3 attempted to put the pill into the Omnicell upon receipt, but the Omnicell was not working at the time. Both LPN10 and LPN3 contacted their Registered Nurse Unit Manager (RN-UM)1 and gave the Oxycodone IR 5 mg pill to RN-UM1 until the Omnicell was working. The DON stated RN-UM1 was interviewed on 09/15/25 and could not locate the Oxycodone IR 5 mg pill. The DON stated RN-UM1 verified she never placed the pill into the Omnicell. The DON stated all medication rooms and carts in the facility were searched without success in locating the pill. The DON stated the police were called, and a report was filed, and RN-UM1 was terminated on 09/15/25. The DON stated the Oxycodone IR 5 mg pill should have been double locked in the medication storage cart until the Omnicell was working. During an interview on 05/15/26 at 9:30 AM, LPN10 revealed one Oxycodone IR 5 mg pill was delivered to the facility on [DATE]. LPN10 stated she signed for the medication and took LPN3 to the medication room for a witness to insert the pill into the Omnicell. LPN10 stated the Omnicell was not working at the time. LPN10 stated she and LPN3 then informed RN-UM1 about the pill being delivered and the Omnicell not working. LPN10 stated RN-UM1 then took the Oxycodone IR 5 mg pill and RN-NM1 stated she would put it into the Omnicell once it was working. LPN10 stated she never heard anything else about the Oxycodone until 09/15/25 when the DON informed her the pill was missing. During an interview on 05/15/26 at 9:45 AM, LPN3 revealed she was with LPN10 on 09/10/25 when they attempted to place the Oxycodone IR 5mg pill into the Omnicell. LPN3 verified the Omnicell was not working, and the pill was immediately given to RN-UM1 who stated she would take possession of the pill until the Omnicell was working. Review of the facility policy titled, Controlled Medication Storage and Disposal, with a revision date of 01/09/2026, revealed Schedule II-V medications (Oxycodone IR is a Schedule II drug with a high potential for abuse with use potentially leading to severe psychological or physical dependence) were to be stored in a double-locked, permanently affixed compartment separate from all other non-narcotic medications. The nurse on duty maintained possession of the key to the controlled medication storage (continued on next page)		

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F 0812 Level of Harm - Minimal harm or potential for actual harm Residents Affected - Some	Procure food from sources approved or considered satisfactory and store, prepare, distribute and serve food in accordance with professional standards. Based on observation, interview, and facility policy review, the facility failed to ensure kitchen staff thoroughly cleaned and air-dried pots and pans prior to storage. In addition, the facility failed to ensure staff with facial hair wore a beard guard for facial hair. These failures increased the risk of foodborne illness by allowing wet pots and pans to be stored before fully drying and by failing to cover facial hair. This had the potential to affect 103 of the 152 residents receiving dietary services. Findings include: During an observation and interview on 05/11/26 at 9:30 AM, [NAME] (C) 1 was observed prepping food for lunch. C1 was observed to have a goatee that was approximately 0.25 &ndash; 0.50 inches in length and was not wearing a beard guard. C1 stated, I didn't think it was long enough to need to be covered. The Regional Food Service Director (RFSD) was present at the time and confirmed C1 was not wearing a beard guard and provided him with one. During an observation and interview on 05/11/26 at 9:35 AM, the Food Service Director (FSD) confirmed two pans, 6 inches by 12 inches by 4 inches; seven pans, 6 inches by 6 inches by 6 inches; five pans, 12 inches by 18 inches by 3 inches cleaned and stacked for use, were still wet. The pans had been stacked while still wet and were not given adequate time to air-dry before stacking. The FSD stated, They [pans] should be dry before they are stacked for use. Being wet increases the risk of contamination. Review of the facility's policy titled, Personnel Adherence to Sanitary Procedures, dated 01/07/26, revealed, Policy: . food services and dietary personnel will be required to adhere to the following sanitary standards. a. Hair nets, caps, beard nets, approved hats, or other effective hair restraints which cover all of the hair, must be worn at all times while on duty in order to keep hair from contacting food and food contact surfaces . Review of the facility's policy titled, Washing Pots and Pans, dated 01/07/26, revealed, . Remove pots and pans from sanitizer and place on drying rack in dish room to air dry. Remove pots and pans in dish room after they dry and move to pan rack in kitchen .		