Printed: 02/22/2025 Form Approved OMB No. 0938-0391

STATEMENT OF DEFICIENCIES AND PLAN OF CORRECTION  (X1) PROVIDER/SUPPLIER/CLIA IDENTIFICATION NUMBER: 085054  NAME OF PROVIDER OR SUPPLIER Cadia Rehabilitation Pike Creek  For information on the nursing home's plan to correct this deficiency, please con-		(X2) MULTIPLE CONSTRUCTION A. Building B. Wing  STREET ADDRESS, CITY, STATE, ZIP CODE 3540 Three Little Bakers Blvd Wilmington, DE 19808	
(X4) ID PREFIX TAG	SUMMARY STATEMENT OF DEFICIENCIES  (Each deficiency must be preceded by full regulatory or LSC identifying information)		
F 0558  Level of Harm - Minimal harm or potential for actual harm  Residents Affected - Few	Reasonably accommodate the need 46448  Based on observations and intervier reach for one (R105) out of five rest Review of R105's clinical record re 11/18/22 - R105 was admitted to the his right side following a stroke, at neck into the windpipe), and deper 2/14/23 11:28 AM - During an interwas hanging off the right side of Ristated, Sometimes. They just chan left it within reach of resident.  2/14/23 3:47 PM - During an obserwas lying off of the right side of his 2/21/23 10:00 AM - The Surveyor ounderneath the bedrail hanging a fice (LPN) confirmed that R105's call be to get down here.	eds and preferences of each resident.  ew, it was determined that the facility facility facility facility facility with a past medical history in racheostomy (small surgical opening the dence on a ventilator (machine that moview with R105, the Surveyor observed 105's bed. The Surveyor asked the resided my linen. The Surveyor picked the vation, R105 was observed lying in between the surveyor process.	cluding paralysis and weakness of nat is made through the front of the oves air in and out of your lungs).  If his call bell not within reach as it ident if this happens often and he call bell up to test the device and the with eyes closed. R105's call bell the right side of his bed take and watching television. E19 try sometimes it just takes me long

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

FORM CMS-2567 (02/99) Previous Versions Obsolete Event ID:

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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 02/23/2023
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F 0582  Level of Harm - Minimal harm or potential for actual harm  Residents Affected - Few	Give residents notice of Medicaid/M 47621  Based on record review, interview (R223) out of three residents review R223's signature or to document R when the resident was discharged facility. Findings include:  Review of R223's clinical record reference of R223's MDS assessment are consistent and reasonable 1/25/23 - R223 was discharged from 2/20/23 - Review of R223's NOMN facility) beneficiary protection notice that she received and understood to 2/21/23 at 9:53 AM - During an interconfirmed that R223's family members 23 confirmed that R223 was not a	full regulatory or LSC identifying information.  Medicare coverage and potential liability and other facility information as indicated wed for the beneficiary protection notification and the Notice of Medicare Part A Services and transfer and the Notice of Medicare Part A Services and transfer and the Notice Part A Services and transfer and Medicare Part A Service	y for services not covered.  ed, it was determined that for one cation, the facility failed to obtain icare Non-Coverage (NOMNC) insferred to an assisted living dent and able to make decisions erred to an assisted living facility.  completed SNF (skilled nursing it sign the statement acknowledging garding the NOMNC process, E23 it date of Medicare Part A services.

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Cadia Rehabilitation Pike Creek		3540 Three Little Bakers Blvd Wilmington, DE 19808		
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(X4) ID PREFIX TAG	PREFIX TAG  SUMMARY STATEMENT OF DEFICIENCIES  (Each deficiency must be preceded by full regulatory or LSC identifying information)		on)	
F 0656	Develop and implement a complete that can be measured.	e care plan that meets all the resident's	needs, with timetables and actions	
Level of Harm - Minimal harm or potential for actual harm	46134			
Residents Affected - Few	Based on record reviews and interviews, it was determined that the facility failed to develop a comprehensive person-centered care plan for two (R102 and R619) out of 35 residents sampled for review of care plans. Findings include:			
	1. Review of R619's clinical record	revealed the following:		
	12/25/22 - R619 was admitted to the	ne facility with multiple diagnoses, inclu	ding respiratory failure and asthma.	
	12/28/22 - A Physician's order was written for Ipratropium-Albuterol Solution 3 ml inhale orally every 6 hours as needed for shortness of breath.			
	12/28/2022 - A Physician's order was written for Continuous Positive Airway Pressure (CPAP) breathing machine use at bedtime.			
	12/29/22 - A Physician's order was asthma.	written for Montelukast Sodium 1 table	et by mouth one time a day for	
	Review of R619's comprehensive prespiratory care for CPAP and trea	person-centered care plan revealed a la tment to be provided to R619.	ack of evidence regarding the	
		terview, E5 (Respiratory Therapist) ack are plan did not contain evidence regar		
	46448			
	2. Review of R102's clinical record	revealed:		
		e facility with a past medical history incl hage, and contractures of the left hip a		
	11/13/22 - R102's comprehensive MDS assessment revealed that he was a totally dependent (restaff performance every time during a 7-day period) in the following activities of daily living (ADL) dressing, eating, toileting, and personal hygiene.			
	Record review lacked evidence of	an ADL care plan for R102.		
	looked through R102's care plan at initiates the care plans? E21 replie	view, E21 (RNAC) was asked for a cop nd stated, I don't see one in here for hir d, The CAA (Care Area Assessment) w AA triggers, if not it's up to nursing to n	n. The Surveyor asked who vill trigger . the resident did not	
	(continued on next page)			

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Cadia Rehabilitation Pike Creek	0.740 74 1444 75 14 75 1		
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F 0656  Level of Harm - Minimal harm or potential for actual harm  Residents Affected - Few	developed to address medical, nurs the comprehensive assessment. A Care plans should include: Service The facility failed to develop and im	y on Care Planning revised on 1/12/23 documented, A comprehensive care plan should be ress medical, nursing, nutritional, and psychosocial needs within 7 days of completion of re assessment. A comprehensive care plan must be prepared by an interdisciplinary team and include: Services furnished to maintain highest practical well-being.  It do develop and implement a comprehensive ADL care plan for R102.  Findings were reviewed with E1 (CNO) and E2 (NHA).	

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F 0657 Level of Harm - Minimal harm or potential for actual harm Residents Affected - Few	Develop the complete care plan with and revised by a team of health processes by a team of healt	thin 7 days of the comprehensive asserblessionals.  ew, it was determined that for one (R9 failed to ensure that the required interd in the care plan meeting. Findings inclusive to the care plan meeting, with a real document of the care plan meeting, with a real document of the care plan meeting and revised by the interval of the care plan meeting. Findings inclusive to the care plan meeting, with a real document of the care plan meeting, with a real document of the care plan meeting. The care plan meeting is called the care plan meeting and the care plan meeting.  The care plan meeting is called the care plan meeting is called the care plan meeting. The care plan meeting is called the care plan meeting in the care plan meeting. The care plan meeting is called the care plan meeting in the care plan meeting.  The care plan meeting is called the care plan meeting. The care plan meeting is called the care plan meeting. The care plan meeting is called the care plan meeting. The care plan meeting is called the care plan meeting. The care plan meeting is called the care plan meeting. The care plan meeting is called the care plan meeting.  The care plan meeting is called the care plan meeting. The care plan meeting is called the care plan meeting. The care plan meeting is called the care plan meeting.  The care plan meeting is called the care plan meeting is called the care plan meeting.  The care plan meeting is called t	ssment; and prepared, reviewed,  7) out of 35 sampled residents for isciplinary team (IDT) members ade:  evision date of 1/12/23, stated that erdisciplinary team (IDT) after each  eleted.  ence of participation by R97's  ce of participation by R97's  was unable to provide evidence ever two Care Conference

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(X4) ID PREFIX TAG	SUMMARY STATEMENT OF DEFICIENCIES (Each deficiency must be preceded by full regulatory or LSC identifying information)		ion)
F 0690 Level of Harm - Minimal harm or potential for actual harm	Provide appropriate care for residents who are continent or incontinent of bowel/bladder, appropriate catheter care, and appropriate care to prevent urinary tract infections.  40163		
Residents Affected - Few	Based on observation, interview and review of the clinical record, it was determined that for one (R108) out of four residents reviewed for urinary catheters/urinary tract infection (UTI), the facility failed to ensure that R108 received appropriate treatment and services to prevent the risk of infection UTI when R108's urinary catheter bag was observed lying on a visibly soiled floor. Findings include:		
	The following was reviewed in R10	8's clinical record:	
	12/17/22 - R108 was admitted to th in the bladder to drain urine).	e facility with a neurogenic bladder an	d required a foley catheter (a tube
	12/21/22 - A Physician's order inclu	ided indwelling catheter care every shi	ft.
	hooked to the left side of R108's be	m observation, R108's foley catheter d ed, approximately one half full of urine oor was observed to be visibly soiled v	and lying on the floor. Beneath
	2/14/23 10:36 AM - During an intervisibly soiled floor.	view, E30 (CNA) confirmed that R108's	s catheter drainage bag was on a
	The facility failed to prevent the risk catheter drainage bag on a visibly s	s for R108 to acquire a bladder infectio soiled floor.	n related to placement of the
	2/23/23 at 2:15 PM - The finding wa (DON) during the exit conference.	as reviewed during the Exit Conference	e with E1 (CNO), E2 (NHA) and E3

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lan to correct this deficiency, please conf	tact the nursing home or the state survey	agency.
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Ensure drugs and biologicals used professional principles; and all drug locked, compartments for controlled 46134  Based on observations and interviewere stored and labeled properly in reviewed. Findings include:  The facility policy on storage of mecontainer, place the date on the container, pla	in the facility are labeled in accordance is and biologicals must be stored in local drugs.  The second second in local drugs is an accordance in local drugs.  The second in local drugs is a second in local drugs.  The second in local drugs is a second in local drugs in local drugs in local drugs is a second in local drugs in local dr	e with currently accepted ked compartments, separately ailed to ensure that medications one out of two medication rooms ed, .When opening a multi-dose was observed:  Ital containing Olopatedine (used for bottle of Keppra (used for seizures) bottle of Omeprazole (used for nding.
	an to correct this deficiency, please configurations and biologicals used professional principles; and all drug locked, compartments for controlled 46134  Based on observations and interviewere stored and labeled properly in reviewed. Findings include:  The facility policy on storage of mecontainer, place the date on the concept of the container, place the date on the concept of	A. Building B. Wing  STREET ADDRESS, CITY, STATE, ZI 3540 Three Little Bakers Blvd Wilmington, DE 19808  an to correct this deficiency, please contact the nursing home or the state survey as  SUMMARY STATEMENT OF DEFICIENCIES (Each deficiency must be preceded by full regulatory or LSC identifying information in the facility are labeled in accordance professional principles; and all drugs and biologicals must be stored in loc locked, compartments for controlled drugs.  46134  Based on observations and interviews, it was determined that the facility fix were stored and labeled properly in two out of five medication carts and in reviewed. Findings include:  The facility policy on storage of medications, last updated 1/31/23, indicate container, place the date on the container.  2/16/23 - During a medication storage review of the first floor the following 2/16/23 10:30 AM- The [NAME] Clay 3 medication cart had one opened by that did not have an open date. E19 (LPN) confirmed the finding.  2/16/23 11:30 AM - The [NAME] Clay 1 medication cart had one opened by that did not have an open date. E19 confirmed the finding.  2/16/23 11:30 AM - The [NAME] Clay medication room had a refrigerated acid reflux) liquid that did not have an open date. E9 (RN) confirmed the finding.

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F 0812  Level of Harm - Minimal harm or potential for actual harm  Residents Affected - Some	Procure food from sources approve in accordance with professional states 36017  Based on observations and intervie handwashing stations were properly The following were observed on 2/1. The cleaning supplies were store access to the handwashing sink.  2. No handwashing sign was present Dietary Director's office.	ed or considered satisfactory and store andards.  ew, it was determined that the facility fa	n, prepare, distribute and serve food siled to ensure that the kitchen's the initial kitchen tour:  the dishwashing room, blocking the handwashing sink outside of the

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(X4) ID PREFIX TAG	SUMMARY STATEMENT OF DEFICIENCIES  (Each deficiency must be preceded by full regulatory or LSC identifying information)		on)	
F 0880	Provide and implement an infection	n prevention and control program.		
Level of Harm - Immediate jeopardy to resident health or	20835			
safety		and reviews of clinical records, facility		
Residents Affected - Few	as indicated, it was determined that the facility failed to maintain an infection prevention and control program to protect the residents against an outbreak of norovirus (a highly contagious infection that can cause a sudden onset of severe vomiting and diarrhea) and they failed to have a system in place to control a contagious disease to residents, staff and visitors. For three (R52, R167 and R417) out of nine sampled residents, the facility failed to actively adhere to contact precautions and hand hygiene among healthcare personnel, residents, and visitors in patient care areas affected by outbreaks of norovirus gastroenteritis, including a process to identify the residents who were symptomatic with signs and symptoms of norovirus as per CDC guidance. The facility's system failure put all residents, including R52, R167 and R417 at immediate jeopardy (IJ) of a serious adverse outcome by not having a method of identifying residents that required contact precautions as it relates to Norovirus, not providing staff the necessary education and PPE (Personal Protective Equipment) to use for contact precautions while delivering resident care and lack of evidence of staff surveillance as per CDC guidance. The IJ was identified on 2/16/23 at 4:11 PM and was abated on 2/20/23 at 10:19 AM. In addition, the facility failed to use infection control precautions for R29 during wound care. Findings include:  The facility policy titled, Standard and Transmission Based Precautions, effective June 2013 and revised 1/27/23 documented, 'Contact Precautions' -applies to all residents infected or colonized with a MDRO (multidrug-resistant organisms or bacteria that resist treatment with more than one antibiotic) in the following situations: presence of acute diarrhea . c. diff (or clostridium difficile, a bacterial overgrow that releases toxins that attack the lining of the intestines) . norovirus . Special Situations: Organisms likely to have spores like Clostridium difficile and some diseases with ongoing transmission like Norovi			
	The CDC's Guideline for the Prevention and Control of Norovirus Gastroenteritis Outbreaks in Healthcare Settings, last updated 2/15/17, documented, . The Summary of Recommendations includes recommendations organized into the following categories:			
	-Patient Cohorting and Isolation Precautions . Avoid exposure to vomitus or diarrhea . During outbreaks, place patients with Norovirus gastroenteritis on Contact Precautions for a minimum of 48 hours after the resolution of symptoms to prevent further exposure of susceptible patients . Consider longer periods of isolation or cohorting precautions for complex medical patients (e.g., those with cardiovascular, autoimmu immunosuppressive, or renal disorders) as they can experience protracted episodes of diarrhea and prolonged viral shedding. Patients with these or other comorbidities have the potential to relapse, and facilities may choose longer periods of isolation based on clinical judgment . Staff who have recovered for recent suspected norovirus infection associated with an outbreak may be best suited to care for symptom patients until the outbreak resolves .			
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F 0880  Level of Harm - Immediate jeopardy to resident health or safety	-Hand Hygiene . Actively promote adherence to hand hygiene among healthcare personnel, patients and visitors in patient care areas affected by outbreaks of norovirus gastroenteritis . During outbreaks, use soap and water for hand hygiene after providing care or having contact with patients suspected or confirmed with norovirus gastroenteritis .		
Residents Affected - Few	-Personal Protective Equipment . If norovirus infection is suspected, adherence to PPE use according to Contact and Standard Precautions is recommended for individuals entering the patient care area (i.e. gowns and gloves upon entry) to reduce the likelihood of exposure to infectious vomitus or fecal material . Use a surgical or procedure mask and eye protection or a full face shield if there is an anticipated risk of splashes to the face during the care of patients, particularly among those who are vomiting .		
	I .	staff, patients, and visitors, including red transmission upon the recognition and	, ,
	-Active Case-Finding . Begin active case-finding when a cluster of acute gastroenteritis cases is detected in the healthcare facility. Use a specified case definition, and implement line lists to track both exposed and symptomatic patients and staff. Collect relevant epidemiological, clinical, and demographic data as well as information on patient location and outcomes .		
	of norovirus gastroenteritis is suspe	Provide timely communication to perso ected and outline what policies and pro s://www.cdc.gov/infectioncontrol/guidel	visions need to be followed to
	According to the facility's completed form entitled Gastroenteritis Data Collection Line Listing for Patients, from 1/31/23 to 2/15/23, 46 residents were documented with gastrointestinal (GI) symptom(s) on both floors of the facility.		
	Observations and clinical record re GI Line Listing revealed:	views of three current residents with re	ecent GI symptoms identified on the
	1a. Review of R167's clinical record	d revealed the following:	
	2/9/23 - R167 was admitted to the	facility.	
	2/13/23 5:47 PM - Review of the Electronic Medication Administration Record (eMAR) revealed that R167 was administered a medication to treat diarrhea and another medication to treat nausea and vomiting. Subsequently, the eMAR documented that these medications were effective in treating the diarrhea, as well as the nausea and vomiting.		
	2/14/23 8:45 AM - A Physician Order was written to chart GI symptoms, which included to monitor for nausea, vomiting, and diarrhea every shift for three days.		
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F 0880  Level of Harm - Immediate jeopardy to resident health or safety	2/15/23 3:30 PM - An interview with R167 revealed that he had nausea yesterday on 2/14/23, but denied nausea, vomiting and/or diarrhea today. An observation of the exterior door into R167's room revealed that there was no signage indicating that R167 was on contact precautions and there was no PPE hanging on his door for staff and visitors to apply prior to entering the room.		
Residents Affected - Few	2/16/23 3:40 PM - An interview with precautions.	n the assigned nurse E8 (LPN) reveale	d that R167 was not on contact
		ad contact precautions implemented (in the aware of the precautions to follow d	
	46134		
	1b. Review of R417's clinical record	d revealed the following:	
	2/8/23 - R417 was admitted to the	facility.	
	2/14/23 3:26 PM - A progress note orders for medications to treat dian	by E24 (NP) stated that R417 was exp hea and nausea and vomiting.	eriencing loose stools and had
	2/15/23 3:45 PM - An interview was done with E6 (CNA) and E7 (CNA). E6 stated that a stomach bug is growing and growing and it is hard to make sure that you don't catch it. E6 and E7 both stated they do not know when residents are positive for a stomach virus and that they ask the Nurses to find out information about residents because the Nurses do not automatically tell them. E6 and E7 added that it is hard to tell who has a stomach virus because some residents are on laxatives and it is hard to know the difference when the residents have loose stools.		
	2/16/23 9:00 AM - A review of the f that R417's last episode of diarrhea	acility's infection control tracking docur a was on 2/15/23 at 2:00 PM.	nent for gastroenteritis documented
		of R417's exterior room door lacked e or staff and visitors to apply prior to en	•
	1c. Review of R52's clinical record	revealed the following:	
	12/23/22 - R52 was admitted to the	facility.	
	2/14/23 3:15 AM - A nurses note in monitored for GI symptoms.	the electronic medical record (EMR) re	evealed that R52 was being
	2/14/23 2:00 PM - A nurses note in the EMR revealed that R52 was experiencing loose stools and a new order was received for a medication to treat R52's diarrhea. R52's onset of diarrhea was noted on the facility's Gastroenteritis: Data Collection Line Listing.		
	(continued on next page)		

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F 0880  Level of Harm - Immediate jeopardy to resident health or safety	2/16/23 11:00 AM - While PPE was present outside of R52's door as the resident was already on enhanced barrier precautions, an observation of R52's exterior door revealed the absence of contact precautions signage to inform staff and visitors to apply the PPE prior to entering the room. According to the facility's Gastroenteritis: Data Collection Line Listing, R52's last episode of diarrhea was on 2/16/23 at 8:30 AM.		
Residents Affected - Few		vations, interviews, review of facility do d reviewed with facility leadership inclu	
	2/16/23 7:02 PM - It was confirmed by Surveyors that the affected three resident rooms had contact precautions signage and PPE in place for staff and visitors on what was required prior to entering the rooms. The facility started staff education that was ongoing around the clock and through an electronic module available to staff. In addition, a CDC sign for norovirus was posted at the receptionist desk educating visitors on what precautions were required during the outbreak.		
	2/16/23 7:30 PM - E2 submitted an	acceptable Abatement Plan signed, da	ated, and timed 2/16/23 at 7:29 PM.
	2/17/23 4:00 PM - E1 was advised completed.	by the Surveyor to send via email evid	ence of staff education once
	2/17/23 5:40 PM - In an email corre PM and stated, continuing to educa	espondence, E2 sent proof of staff educate staff.	cation status as of 2/17/23 at 5:30
	2/20/23 10:19 AM - The date and ti	ime E2 stated that staff in-service was	completed.
	40264		
	The facility failed to provide evid staff for GI signs and symptoms.	ence of an infection control surveillance	e program that included monitoring
	The CDC's Guideline for the Prevention and Control of Norovirus Gastroenteritis Outbreaks in Healthcare Settings, last updated on 2/15/17, documented . The Summary of Recommendations includes recommendations organized into the following categories:		
	-Staff Leave and Policy . Exclude ill personnel from work for a minimum of 48 hours after the resolution of symptoms. Once personnel return to work, the importance of performing frequent hand hygiene should be reinforced, especially before and after each patient contact .		
	2/14/23 12:17 PM - An interview with E9 (LPN) stated that a GI illness started on the second floor last weekend and then spread to the first floor residents. E9 stated that she believed a random test was done, but she was not sure if the residents were tested specifically for Norovirus.		
	(continued on next page)		

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 02/23/2023
NAME OF PROVIDER OR SUPPLIER		STREET ADDRESS, CITY, STATE, ZIP CODE	
Cadia Rehabilitation Pike Creek		3540 Three Little Bakers Blvd Wilmington, DE 19808	
For information on the nursing home's	plan to correct this deficiency, please con	tact 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 0880  Level of Harm - Immediate jeopardy to resident health or safety	2/15/23 3:00 PM - E3 (IC/DON) provided the survey team with copies of the correspondence from the State of Delaware Department of Public Health (DPH). This included the Gastroenteritis Data Collection Line List. DPH recommended 2-3 resident stool samples for testing. E3 also stated that the official start of the outbreak was 2/8/23 when the facility received notification of a positive test for norovirus.		
Residents Affected - Few	2/16/23 10:35 AM - A joint interview with E1 (CNO), E2 (NHA) and E3 revealed that DPH sent an email for the facility to call DPH to go over the results and recommendations when the positive norovirus test came back on 2/8/23. According to E1 and E3, the recommendations included to quarantine (resident) cases in their room .if .any more staff please exclude them until .(diarrhea) symptoms have resolved .isolating precautions .cohorting or exclusion for staff .		
	2/16/23 10:45 AM - An interview with E3 confirmed that an infection control line listing was not initiated for staff. It was further revealed there were sporadic staff cases based on who called out sick. E3 stated that symptomatic staff could not come back to work until they were without nausea, vomiting, diarrhea and/or fever for 48 hours. E3 stated that she'll start creating the staff line list using the DPH template.		
	47114		
	3. Review of the facility policy and procedure, revised 6/2/2021, titled Infection Control Hand Hygiene included, .it is the policy of Cadia Healthcare to help control the spread of infection through hand hygiene, wash hands thoroughly, using rigorous scrubbing action for at least 20 seconds.		
	Review of When and How to Perform Hand Hygiene included .after touching a patient or the patient's immediate environment, after contact with blood, body fluids or a contaminated surface. https://www.cdc.gov/handhygiene/providers/index.html.		
	2/20/23 9:55 AM - During an observation, R29's breakfast tray and a urinal containing urine were on the bedside table. E14 (LPN) placed dressing supplies on the unclean bedside table and then picked up R29's breakfast tray and left the room. E14 then picked up the urinal from the bedside table and emptied it in the bathroom and washed her hands for five seconds. E14 placed a blue pad on the bedside table to create a clean field, then picked up the dressing supplies that had been placed on a dirty bedside table, and put them on the blue pad causing contamination of the clean field. E14 then put on clean gloves and opened the treatment supplies that had been contaminated causing the nurses gloves to no longer be clean. The nurse proceeded to complete the dressing change with gloved hands that were contaminated.		
	2/21/23 11:00 AM - The Surveyor asked E1 (CNO) for a policy and procedure for wound care, and E1 replied, No, we don't have a policy and procedure for wound care.		
	2/21/23 2:11 PM - During an interview with E14 she stated, I put the dressings on the table and put the dressings on top of the pad.		
	2/23/23 - Findings were reviewed with E1 (CNO), E2 (NHA) and E3 (DON) during the exit conference beginning at 2:15 PM.		