

STATEMENT OF DEFICIENCIES AND PLAN OF CORRECTION	(X1) PROVIDER/SUPPLIER/CLIA IDENTIFICATION NUMBER: 355093	(X2) MULTIPLE CONSTRUCTION A. Building B. Wing	(X3) DATE SURVEY COMPLETED 09/18/2024
NAME OF PROVIDER OR SUPPLIER Good Samaritan Society - Bottineau		STREET ADDRESS, CITY, STATE, ZIP CODE 725 E 10th St Bottineau, ND 58318	

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)
<p>F 0641</p> <p>Level of Harm - Minimal harm or potential for actual harm</p> <p>Residents Affected - Few</p>	<p>Ensure each resident receives an accurate assessment.</p> <p>**NOTE- TERMS IN BRACKETS HAVE BEEN EDITED TO PROTECT CONFIDENTIALITY** 46259</p> <p>Based on record review, review of the Long-Term Care Facility Resident Assessment Instrument (RAI) 3.0 User's Manual (Version 1.18.11), and staff interview, the facility failed to ensure accurate coding of the Minimum Data Set (MDS) for 3 of 13 sampled residents (Resident #11, #12, and #26). Failure to accurately complete the MDS does not allow each resident's assessment to reflect their current status/needs and may affect the accurate development of a comprehensive care plan and the care provided to the residents.</p> <p>Findings include:</p> <p>SECTION K: SWALLOWING/NUTRITIONAL STATUS</p> <p>The Long-Term Care Facility RAI 3.0 User's Manual, revised October 2023, page K-11, stated, . Coding Instructions Check all that apply. K0520B, feeding tube - nasogastric or abdominal (PEG).</p> <p>- Review of Resident #26's medical record occurred on all days of survey. The quarterly MDS, dated [DATE], identified a feeding tube. Review of the physician's orders for the assessment period lacked evidence of a feeding tube.</p> <p>SECTION M: SKIN CONDITIONS</p> <p>The Long-Term Care Facility RAI 3.0 User's Manual, revised October 2023, pages M-5 and M-12/13, stated, [M-5] . Coding Instructions Code based on the presence of any pressure ulcer/injury in the past 7 days. Code 1, yes: if the resident had any pressure ulcer/injury (Stage 1, 2, 3, 4, or unstageable) in the 7-day look-back period. [M-12/13] . Stage 2 Pressure Ulcer Partial thickness loss of dermis presenting as a shallow open ulcer with a red or pink wound bed, without slough. May also present an intact or open/ruptured blister. Coding Instructions for . Enter the number of pressure ulcers that are currently present and whose deepest anatomical stage is Stage 2. Enter the number of these 2 pressure ulcers that were first noted at the time of admission/entry AND-for residents who are reentering the facility after a hospital stay, enter the number of Stage 2 pressure ulcers that were acquired during the hospitalization .</p> <p>- Review of Resident #11's medical record occurred on all days of survey. The record showed the resident returned from the hospital on 08/08/24.</p> <p>Review of a nurses' note 08/09/24, stated, . Blister to the left heel [sic] is just beginning, .</p> <p>(continued on next page)</p>

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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<p>F 0641</p> <p>Level of Harm - Minimal harm or potential for actual harm</p> <p>Residents Affected - Few</p>	<p>The quarterly MDS, dated [DATE], failed to identify an unhealed pressure ulcer.</p> <p>SECTION N: MEDICATIONS</p> <p>The Long-Term Care Facility RAI 3.0 User's Manual, revised October 2023, pages N-13, stated, . Review the resident's medication administration records to determine if the resident received an antipsychotic medication . Coding Instructions . Code 0, no: if antipsychotics were not received . Code 1, yes: if antipsychotics were received on a routine basis only .</p> <p>- Review of Resident #12's medical record occurred on all days of survey. A physician's order dated 04/12/24 stated, Seroquel [antipsychotic medication] Oral Tablet 100 MG [milligrams] Give 100 mg by mouth two times a day .</p> <p>The quarterly MDS, dated [DATE], failed to reflect the resident's routine use of an antipsychotic medication.</p> <p>During an interview on 09/08/24 at 12:03 p.m., an administrative staff member (#1) confirmed Resident #11, #12, and #26's MDS assessments were coded incorrectly.</p> <p>13101</p>

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<p>F 0658</p> <p>Level of Harm - Minimal harm or potential for actual harm</p> <p>Residents Affected - Few</p>	<p>Ensure services provided by the nursing facility meet professional standards of quality.</p> <p>39211</p> <p>Based on observation, record review, facility policy review, and staff interview, the facility failed to follow professional standards of practice for 2 of 2 sampled residents (Resident #5 and #11) observed with an indwelling catheter. Failure to obtain physician orders for an indwelling catheter, catheter care, and/or maintenance of the catheter may result in an adverse consequence for residents.</p> <p>Findings include:</p> <p>Review of the facility policy titled Catheter: Care, Insertion & Removal, Drainage Bags, Irrigation, Specimen occurred on 09/18/24. This policy, dated 07/30/24, stated. Catheters will be inserted only with a physician's order and will include the type of catheter, size and balloon capacity for indwelling catheter. Indwelling or retention catheters are changed only when necessary and are connected to a closed drainage system. All closed collection systems that become contaminated by inappropriate technique, leaks or other means are immediately replaced. They are also changed when a new catheter is inserted and at other times only when necessary due to encrustations or according to physician's orders.</p> <p>Review of the facility policy titled Physician/Practitioner Orders occurred on 09/18/24. This policy, dated 04/01/24, stated, . The admitting orders are intended to provide guidance on appropriate resident care . Required orders on admission: . Treatments: a treatment order will include the supporting reason (diagnosis/problem) .</p> <p>- Review of Resident #5's medical record occurred on all days of survey. The physician orders, dated 06/26/24, included placement of a foley (indwelling) catheter. Observation on all days of survey showed Resident #5 with an indwelling urinary catheter. The physician orders lacked instructions for catheter changes and/or care and maintenance of the indwelling catheter.</p> <p>During an interview on the morning of 09/18/24, an administrative staff member (#7) confirmed Resident #5's physician orders lacked instructions for the care and management for the indwelling catheter.</p> <p>13101</p> <p>- Review of Resident #11's medical record occurred on all day s of survey. Observations on September 16 and 17, 2024 showed Resident #11 with an indwelling urinary catheter. The physician orders lacked an order for the catheter, instruction for catheter changes and/or care and maintenance of the indwelling catheter.</p> <p>On the morning of 09/18/24, an administrative staff member (#7) confirmed the facility staff failed to transcribe Resident #11's physician orders for the indwelling catheter.</p>		

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<p>F 0759</p> <p>Level of Harm - Minimal harm or potential for actual harm</p> <p>Residents Affected - Few</p>	<p>Ensure medication error rates are not 5 percent or greater.</p> <p>13101</p> <p>Based on observation, review of manufacturer's instructions, and staff interview, the facility failed to ensure a medication error rate of less than five percent for 3 of 5 residents (Resident #9, #17, and #25) observed during medication administration. Four medication errors occurred during staff administration of 25 medications, resulting in a 16 percent error rate. Failure to accurately prepare and administer medication may result in residents receiving an ineffective dose and/or experiencing adverse reactions.</p> <p>Findings include:</p> <p>Review of the manufacturer's package insert for Fiasp (fast acting insulin) FlexTouch Pen, dated October 2019, stated, [Preparing Pen] . Step 4: Push the capped needle straight onto the Pen and twist the needle on until it is tight. Step 5: Pull off the needle cap. Do not throw it away. Step 6: Pull off the inner needle cap and throw it away. [Priming Pen] Step 7 Turn the dose selector to select 2 units . Step 9: Hold the pen with the needle pointing up. Press and hold in the dose button until the dose counter shows '0'. The '0' must line up with the dose pointer. A drop of insulin should be seen at the needle tip . [Select dose] . [Giving injection] . Step 13 Press and hold down the dose button until the dose counter shows '0' . Keep the needle in your skin after the dose counter has returned to '0' and slowly count to 6. When the does counter returns to '0', you will not get your full dose until 6 seconds later.</p> <p>Observations of Fiasp insulin administration for Resident #25:</p> <p>- On 09/17/24 at 4:42 p.m. showed a nurse (#3) primed the insulin pen with the needle cap on and pointed down. The nurse (#3) failed to prime the pen with the cap off and the needle pointed up.</p> <p>- On 09/18/24 at 11:49 a.m. showed a nurse (#4) turned the dose selector to 6 units as ordered, attached the needle, and without priming the insulin pen administered the insulin, and removed the pen immediately from the resident's skin. The nurse (#4) failed to correctly attach the needle, prime the pen, and keep the needle in the skin for the recommended amount of time.</p> <p>Manufacturer's directions for polyethylene glycol (a powdered laxative) located on the medication bottle, stated, Directions . fill to the top of the bottle cap which will provide the correct dose (17 g [grams]), stir and dissolve in any 4 to 8 ounces of beverage . then drink .</p> <p>Observations on 09/18/24:</p> <p>- At 7:49 a.m., showed a medication aide (MA) (#5) placed 17 g of polyethylene glycol in a cup, added two to three ounces of water, and stirred. After Resident #9 drank the liquid, residue from the medication remained on the inside of the cup.</p> <p>(continued on next page)</p>		

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<p>F 0759</p> <p>Level of Harm - Minimal harm or potential for actual harm</p> <p>Residents Affected - Few</p>	<p>- At 8:02 a.m., showed the MA (#5) placed 17 g of polyethylene glycol in a cup, added two to three ounces of water, and stirred. The MA (#5) held the cup so Resident #17 could drink the medication through a straw. The resident left half to one ounce of medication, not fully dissolved, in the cup. The MA (#5) failed to use the recommended amount of water to dissolve the ordered dose of polyethylene glycol.</p> <p>During an interview on 09/18/24 at 12:13 p.m., three administrative staff members (#6, #7, and #8) confirmed facility staff failed to administer medications according to manufacturer's recommendations.</p>		

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<p>F 0880</p> <p>Level of Harm - Minimal harm or potential for actual harm</p> <p>Residents Affected - Few</p>	<p>Provide and implement an infection prevention and control program.</p> <p>39211</p> <p>Based on observation, record review, facility policy review, and professional reference, the facility failed to follow standards of infection control for 2 of 2 sampled residents (Resident #5, and #11) observed during catheter cares. Failure to follow infection control practices during resident cares related to hand hygiene, glove use, emptying urine container, and enhanced barrier precautions (EBP), has the potential to spread infection throughout the facility.</p> <p>Findings include:</p> <p>Review of facility policy titled Catheter: Care, Insertion & Removal, Drainage Bags, Irrigation, Specimen occurred on 09/18/24. This policy, dated 07/30/24, stated. Catheter Drainage Bag Emptying clinical skill check list. perform hand hygiene. Apply gloves. When emptying the catheter bag, place a fluid-impermeable pad beneath the measuring container and avoid placing the measuring container on the floor. Wash and dry measuring container. Remove gloves, perform hand hygiene, .</p> <p>Review of facility policy titled Standard and Transmission-Based Precautions occurred on 09/18/24. This policy, dated 04/02/24, stated. Enhanced barrier precautions . refer to the use of gown and gloves during high-contact resident care activities . are needed for residents with . Indwelling Medical devices (.indwelling urinary catheters .). High-Contact Resident Care Activities include: Transfers, . while assisting with transfers and mobility, .</p> <p>Kozier & Erb's Fundamentals of Nursing: Concepts, Process, and Practice, 11th ed., Pearson Education, Incl, New Jersey, 2021, pages 686-687, stated, Skill 31.2 Applying and Removing Personal Protective Equipment (Gloves, Gown, Mask, Eyewear) . 3. Apply a clean gown. Overlap the gown at the back as much as possible, and fasten the waist ties or belt. 6. Apply clean gloves.</p> <p>- Review of Resident #5's medical record occurred on all days of survey. The care plan indicated EBP related to an indwelling urinary catheter. A supply cart and signage for EBP observed inside the resident's room.</p> <p>Observation on 09/16/24 at 1:17 p.m. showed a certified nurse aide (CNA) (#2) entered Resident #5's room. The CNA positioned Resident #5 next to the bed, reached under the resident's right arm, assisted the resident to a standing position from the wheelchair, and pivot transferred the resident onto the bed. The CNA lifted the resident's legs onto the bed and hung the urine collection bag onto the bed frame. The CNA (#2) donned a gown and gloves, emptied the urine collection bag into a measuring container, discarded the urine into the toilet, held the contaminated container under the sink faucet, of a shared bathroom, obtained water, rinsed, and emptied the contents of the container into the toilet.</p> <p>The CNA (#2) failed to don a gown or gloves before assisting with a transfer, failed to perform hand hygiene prior to donning PPE, and held a contaminated urine container in the sink of a shared bathroom.</p> <p>13101</p> <p>(continued on next page)</p>		

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<p>F 0880</p> <p>Level of Harm - Minimal harm or potential for actual harm</p> <p>Residents Affected - Few</p>	<p>- Review of Resident #11's medical record occurred on all days of survey. The care plan indicated EBP related to an indwelling urinary catheter. Signage for EBP was located on the hallway door frame and a supply cart was in the resident's room.</p> <p>Observation on 09/17/24 at 9:07 a.m., showed a CNA (#9) assisted Resident #11 with a brief change. The CNA (#9) wore a gown and gloves, but failed to tie the gown at the waist. The untied gown kept falling into the workspace, and the CNA (#9) stopped cares, lifted her arms, and shrugged both shoulders, to get the gown out of the workspace. The CNA (#9) changed gloves without hand hygiene, obtained a measuring container and without a barrier placed the container on the floor. The CNA (#9) emptied most of the urine into the container, her gown falling forward, stood emptied the urine into the toilet, the gown wrapped around the front half of the toilet, repeated the process a second time to fully empty the urine collection bag, and without rinsing/cleansing the container put it away. The CNA (#9) removed her gown and gloves and without performing hand hygiene left the resident's room to obtain assistance. The CNAs (#9 and #10) entered the resident's room, CNA (#10) entered the bathroom, donned gloves, and returned to the door to don a gown. CNA (#9) donned a gown, tied it only at the neckline and entered the bathroom to don gloves. After the CNAs (#9 and #10) transferred Resident #11, CNA (#10) removed the gown/gloves and washed her hands. CNA (#9) changed gloves and without performing hand hygiene assisted Resident #11 with grooming and mouth care. CNA (#10) collected the garbage, started to disinfect the lift without gloves, stopped donned gloves, disinfected the lift, removed her gloves, and exited the room without performing hand hygiene. The CNAs failed to use PPE appropriately, one CNA (#9) failed to tie the gown at the waist and the other CNA (#10) failed to don the gown and then the gloves.</p>		