Printed: 12/22/2024 Form Approved OMB No. 0938-0391

STATEMENT OF DEFICIENCIES AND PLAN OF CORRECTION	(X1) PROVIDER/SUPPLIER/CLIA IDENTIFICATION NUMBER: 505296	(X2) MULTIPLE CONSTRUCTION  A. Building  B. Wing	(X3) DATE SURVEY COMPLETED 05/31/2023
NAME OF PROVIDER OR SUPPLIER St Francis of Bellingham		STREET ADDRESS, CITY, STATE, ZIP CODE 3121 Squalicum Parkway Bellingham, WA 98225	
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 0686 Level of Harm - Actual harm Residents Affected - Few			evelop a plan of care, and ) development for 1 of 1 resident morbidities with an increased risk re competency on the correct use of uency energy to the bone), prevent the bone stimulator. This failure Pl to their left foot from a medical  at Assessment Instrument manual, erlying tissue, usually over a bony in combination with shear. The be painful. A stage III PU is defined endon or muscle is not exposed. The depth of tissue loss.  dated January 2020, showed PU oen). Medical device related PIs apeutic purposes. The resultant PI d be staged using the staging on of medical device-related  ally contraindicated).  down.

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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			NO. 0936-0391	
STATEMENT OF DEFICIENCIES AND PLAN OF CORRECTION	(XI) PROVIDER/SUPPLIER/CLIA IDENTIFICATION NUMBER: 505296	(X2) MULTIPLE CONSTRUCTION  A. Building  B. Wing	(X3) DATE SURVEY COMPLETED 05/31/2023	
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(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 - Actual harm Residents Affected - Few				
	(continued on next page)			

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STATEMENT OF DEFICIENCIES AND PLAN OF CORRECTION	(XI) PROVIDER/SUPPLIER/CLIA IDENTIFICATION NUMBER: 505296	(X2) MULTIPLE CONSTRUCTION  A. Building  B. Wing	(X3) DATE SURVEY COMPLETED 05/31/2023
NAME OF PROVIDER OR SUPPLIE	ER	STREET ADDRESS, CITY, STATE, ZIP CODE	
St Francis of Bellingham		3121 Squalicum Parkway Bellingham, WA 98225	
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(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 - Actual harm		ale (assessment tool to predict PU risk) PU risk was a score between 15 to 18	
Residents Affected - Few	Nursing Assistant (CNA), notified S	ote, dated 05/15/2023 at 7:09 AM, show staff D, Licensed Practical Nurse (LPN) measuring 0.5 centimeters (cm) by 0.5 e stimulator was on.	, there was an open area on the
	Review of Resident 1's skin and wound evaluation, dated 05/16/2023 at 10:11 AM, showed a new left later: (side) malleolus (ankle) wound that was a medical device related PI, acquired in-house at a stage III with fu thickness skin loss measuring 1.2 cm in length by 0.9 cm in width, by 0.1 cm depth. The wound bed had 50 granulation (new skin) tissue and 50% slough (non-viable tissue).  Review of a progress note on 05/16/2023 at 3:49 PM, showed the factors contributing to the event/incident were Resident 1 had a bone stimulator on their left ankle to support healing after previous fracture. The not stated that the strap of the stimulator went over the open area and the pressure and friction from the device likely caused the skin breakdown. Resident 1had edema (swelling) to the foot and ankle that would contribute to higher risk of skin breakdown. The analysis was the facility determined the wound was a device related PI from the bone stimulator they wore for three hours each night shift. The device centered over the lateral malleolus with the strap wrapping around the ankle. The device vibrated and caused the strap to pus and rub on edematous (swollen) skin.		
	Assistant, documented the residen The wound team saw yesterday an stimulator. However at this time ref can be discontinued. At this time th adjusting diuretics and the patient's still fairly swollen. The patient is go continue to monitor this wound. The that she broke. The patient has a s area was approximately 1.2 cm and orders for Medihoney (a type of wo	ord, dated 05/15/2023 at 7:40 AM, Staft was seen today for evaluation of the vid felt like it was getting worse and requiered back from the patient's orthopedial patient has a small wound in the creasing the patient has a small wound in the creasing the stage of the softer with less pitting eddod about wearing her compression socie patient continues to complain of foot mall open area of the skin where her bid is small. There is some slough present und gel). There is no surrounding signate regimen and monitor daily for any other	wound on the crease of her foot. Dested a hold on the patient's bone of and they have indicated that it ase of her ankle. We have been area (swelling) however the feet are less however at this time we need to pain on the left side and the foot one stimulator was placed. The not and as such yesterday I put in a of cellulitis or significant concern.
	(continued on next page)		

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For information on the nursing home's	nformation on the nursing home's plan to correct this deficiency, please contact the nursing home or the state survey agency.		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 - Actual harm Residents Affected - Few			ed and they felt the resident's didentified that the facility had no ere was no area for skin inspection literature on the bone stimulator ter the injury occurred. Staff B said e on 01/19/2023. Staff B said they re unaware there was a full staff called the orthopedic surgeon's 2023. Staff B said the facility A and Staff B stated there was no they were working on that now.  I was lying in bed. The in that measured approximately 0.9 ithelial tissue. The resident stated ated the wound looked better than se especially with the stimulator did not vibrate at all, it was staff D said that Staff H, NAC, found is compression stockings. Staff D say would place the bone stimulator ent got up in the morning. Staff D ent. Staff D stated they were staff F, LPN/Staff Development ated when they first observed the cm by 0.5 cm. Staff D stated they Staff D said they received no ing it.  In or recent PU prevention education.  at six of twenty-four NACs had a documented to have completed had just talked to Staff A about the on 05/14/2023. Staff E said they	

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