LICENSURE UNIT

PRINTED: 08/25/2015 FORM APPROVED

JEP 22 2015

Nebraska DHHS Licensure Unit (X3) DATE SURVEY (X2) MULTIPLE CONSTRUCTION (X1) PROVIDER/SUPPLIER/CLIA STATEMENT OF DEFICIENCIES COMPLETED AND PLAN OF CORRECTION IDENTIFICATION NUMBER: A. BUILDING: C B. WING_ 08/06/2015 HC001

NAME OF PROVIDER OR SUPPLIER

STREET ADDRESS, CITY, STATE, ZIP CODE

NAME OF I	THO VIDENCE OF CO.	T MISSION	TATE, ZIP CODE					
BELLEVUE HEALTH CENTER BELLEVUE, NE 68005								
(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) COMPLETE DATE				
G 150	7-006.06 Patient Care and Treatment Each health clinic must establish and implement written policies and procedures that encompass all care and treatment provided to patients. The policies and procedures are consistent with prevailing professional standards, delineate the scope of services provided in the health clinic and encompass aspects to protect the health and safety of patients. This Standard is not met as evidenced by: Based on observation, staff interview and policy review; the facility failed to have a policy in place to consistently identify tissue specimen(s) removed during the abortion procedure (extraction of fetal tissue from the uterus) that were stored in the freezer. Five of Five specimens in the freezer were not consistently identified. This procedure had the potential to effect any tissue specimen(s) stored by the facility. Findings are:	G 150						
	A. During the facility tour on 8/4/15 from 12:00 PM to 1: 20 PM; the freezer (which had been identified for storage of tissue specimens) was observed to have five tissue specimen(s) with the following identification: -Specimen 1 A tissue specimen wrapped in a chux (a water impermeable pad) placed in a plastic bag identified with initials and a date written with a magic marker; -Specimen 2 A tissue specimen wrapped in a chux placed in a plastic bag identified with a first initial, last name and a patient number written with a magic marker; -Specimen 3 A tissue specimen wrapped in a chux placed in a plastic bag identified with the		G 150 7-006.06 A. Specimen 1-5 were all disposed of in red biohazard bags and in biohazard box in garage. The DON will be responsible for Checking the freezer monthly.	08/04/1				

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

words 'room 1 specimen' and the date written

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FORM APPROVED Nebraska DHHS Licensure Unit (X3) DATE SURVEY (X2) MULTIPLE CONSTRUCTION STATEMENT OF DEFICIENCIES (X1) PROVIDER/SUPPLIER/CLIA COMPLETED **IDENTIFICATION NUMBER:** AND PLAN OF CORRECTION A. BUILDING: C B. WING 08/06/2015 HC001 STREET ADDRESS, CITY, STATE, ZIP CODE NAME OF PROVIDER OR SUPPLIER 1002 WEST MISSION **BELLEVUE HEALTH CENTER** BELLEVUE, NE 68005 PROVIDER'S PLAN OF CORRECTION SUMMARY STATEMENT OF DEFICIENCIES (X5)(X4) ID COMPLETE (EACH CORRECTIVE ACTION SHOULD BE (EACH DEFICIENCY MUST BE PRECEDED BY FULL PREFIX PREFIX CROSS-REFERENCED TO THE APPROPRIATE DATE REGULATORY OR LSC IDENTIFYING INFORMATION) TAG TAG DEFICIENCY) G 150 G 150 Continued From page 1 with a magic marker; - Specimen 4-- A tissue specimen wrapped in a chux placed in a plastic bag identified with a first name, last name and the date written with a magic marker; and - Specimen 5 -- A tissue specimen wrapped in a chux placed in a plastic bag without any identifying information such as a name, number or date written on the bag. G 150 7-006.06 B-D B. An interview with the Director of Nurses [DON] The policy for Products of 08/07/15 during the tour on 8/4/15 from 12:00 PM to 1:20 Conception has been updated. PM revealed, that there needed to be a better system for identification of the specimens in the freezer. The DON identified that there was not a See Attachments 1-3 specific policy or procedure in place for labeling and identifying specimens placed in freezer for hold. C. Per a written response as part of the physician interview dated 8/13/15, the sole exception to the disposal of tissue via protocol for the certified medical waste disposal company pick up would be: "a) A request from a referring provider to have the tissue forwarded to a laboratory for further diagnostic study. b) A request from a law enforcement agency or jurisdiction to have the tissue surrendered to an agent for evidence. c) A request from the patient to have the tissue released to a licensed funeral director or a agent to prepare the fetus for cremation or burial." G 150 7-006.06 D D. Review of the facility Policy and Procedure Amendment to Dr. Carhart's Manual. Section 8 -Procedure Manual:

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Identification of Products of Conception (POC)

"...If the POC was a result of rape, it will be put in

sterile specimen cup, labeled and placed in the

revealed the following information:

6899

Response.

See Attachment 24

Nebraska DHHS Licensure Unit (X3) DATE SURVEY (X1) PROVIDER/SUPPLIER/CLIA STATEMENT OF DEFICIENCIES (X2) MULTIPLE CONSTRUCTION COMPLETED AND PLAN OF CORRECTION **IDENTIFICATION NUMBER:** A. BUILDING: C B WING 08/06/2015 HC001 STREET ADDRESS, CITY, STATE, ZIP CODE NAME OF PROVIDER OR SUPPLIER 1002 WEST MISSION BELLEVUE HEALTH CENTER BELLEVUE, NE 68005 PROVIDER'S PLAN OF CORRECTION SUMMARY STATEMENT OF DEFICIENCIES (X4) ID (EACH CORRECTIVE ACTION SHOULD BE COMPLETE (EACH DEFICIENCY MUST BE PRECEDED BY FULL **PREFIX** PRÉFIX CROSS-REFERENCED TO THE APPROPRIATE DATE REGULATORY OR LSC IDENTIFYING INFORMATION) TAG TAG DEFICIENCY) G 150 G 150 Continued From page 2 freezer." The Policy and Procedure Manual lacked any further protocol regarding the identification and management of frozen tissue specimens. G 410 7-006.09E Storage of Drugs/Devices/Biologicals G 410 All drugs, devices, and biologicals must be stored in secured areas and stored in accordance with the manufacturer's, distributor's, packager's, or dispensing pharmacist 's instructions for temperature, light, humidity, and other storage instructions. Only authorized personnel, designated by policy and procedure of the health clinic as responsible for administration, provision, or dispensing, must have access to drugs, devices, and biologicals. The supply of drugs, devices, and biologicals must be protected and restricted to use for legally authorized purposes and must be checked on a regular basis to ensure expired, mislabeled, unlabeled, or unusable products are not available for patient use. This Standard is not met as evidenced by: Based on observation and staff interview; the facility failed to ensure that expired biologicals were not available for patient use. Two of Two exam room cupboards contained boxes of Lamicel Osmotic Cervical Dilators and Laminaria Tents [a thin rod of dried kelp that is placed into the cervix (the "neck" of the uterus) to soften and dilate (open) the cervix prior to the abortion procedure (extraction of fetal tissue from the uterus)]. [Each box contained 20-24 Cervical Dilators in individualized pouches.] This had the potential to effect all patients requiring the use of

this product for an abortion procedure.

Nebraska DHHS Licensure Unit (X3) DATE SURVEY STATEMENT OF DEFICIENCIES (X2) MULTIPLE CONSTRUCTION (X1) PROVIDER/SUPPLIER/CLIA COMPLETED **IDENTIFICATION NUMBER:** AND PLAN OF CORRECTION A. BUILDING: C B. WING 08/06/2015 HC001 STREET ADDRESS, CITY, STATE, ZIP CODE NAME OF PROVIDER OR SUPPLIER 1002 WEST MISSION BELLEVUE HEALTH CENTER BELLEVUE, NE 68005 PROVIDER'S PLAN OF CORRECTION (X5) COMPLETE SUMMARY STATEMENT OF DEFICIENCIES (X4) ID (EACH CORRECTIVE ACTION SHOULD BE (EACH DEFICIENCY MUST BE PRECEDED BY FULL PREFIX PREFIX DATE CROSS-REFERENCED TO THE APPROPRIATE REGULATORY OR LSC IDENTIFYING INFORMATION) TAG TAG DEFICIENCY) G 410 G 410 Continued From page 3 Findings are: G 410 7-006.09E A1-2 A. Observations made on the facility tour 8/14/15 All expired supplies were 08/04/15 from 12:00 PM to 1:20 PM revealed the following: disposed of properly. The DON will check for 1) Exam Room 1 had a cupboard that contained expiration dates monthly multiple boxes of Lamicel Osmotic Cervical Dilators and Laminaria Tents. The following boxes of Laminaria/Lamicel Osmotic Cervical Dilators were outdated: -5 boxes of 4 mm (millimeter long) / 70 mm (millimeter diameter) Laminaria Tents which were outdated 9/2014 and 1 box that was outdated 11/2013; -1 box of 2 mm / extra small Laminaria Tents with an outdate of 1/2014 and 1 box that outdated 7/2015; -1 box of 6 mm / 70 mm Laminaria Tents with an outdate of 12/2011; 4 boxes that outdated 7/2014; and 1 box that outdated 7/2015; -1 box of 3 mm / (no other mm listing on box related to diameter) of Lamicel Osmotic Cervical Dilators with an outdate of 8/2005; 1 box with an outdate of 10/2005; 1 box with an outdate of 5/2008 and 1 box with an outdate of 7/2008; and -2 boxes of 5 mm / (no other mm listing on box related to diameter) of Lamicel Osmotic Cervical Dilators with an outdate of 7/2008 and 1 box outdated 12/2003. 2) Exam Room 2 had a cupboard that contained multiple boxes of Laminaria Tent. The following boxes of Laminaria were outdated: -1 boxes of 4 mm (millimeter long) / 70 mm (millimeter diameter) Laminaria Tents with an outdate of 11/2013; -1 box of 2 mm / extra small Laminaria Tents with an outdate of 3/2014; and

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-1 box of 8 mm / 70 mm Laminaria Tents with an

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STATEMENT OF DEFICIENCIES AND PLAN OF CORRECTION		(X1) PROVIDER/SUPPLIER/CLIA IDENTIFICATION NUMBER:	(X2) MULTIPLE CONSTRUCTION		(X3) DATE SURVEY COMPLETED					
AND FLAN OF CORRECTION			A. BUILDING:		C					
		HC001	B. WING		3	6/2015				
NAME OF PROVIDER OR SUPPLIER STREET ADDRESS, CITY, STATE, ZIP CODE										
BELLEVUE HEALTH CENTER 1002 WEST MISSION BELLEVUE, NE 68005										
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G 410	Continued From page 4		G 410							
	outdate of 2/2015. B. Staff interview (during the tour on 8/4/15 from 12:00 PM to 1:20 PM) with the Director of Nurses			G 410 7-006.09E New policies written for supplies.		08/07/15				
		ed, "I do monthly checks to								
	check for expired medications, but didn't realize that those had outdated."			See Attachments 4-5						
G 530	7-006.15B Equipme	ent, Fixtures, and Furnishings	G 530							
	equipment, fixtures, and in good repair. 7-006.15B1 The fimplement a process preventative maintefurnishings to ensur	and maintain all and furnishings clean, safe acility must establish and as designed for routine and anance of equipment and a te that such equipment the								
	Based on observation facility failed to: 1) est maintenance process pieces of medical emachines - medical waves to produce in inside the body; 2 doused only for monitor 1 cautery machine - and repair tissue; 2 which has a tube that removed tissue or flimplement prevental sampled pieces of mautoclaves - a machinstruments in betwee practice has the potential process.	on and staff interview; the stablish preventative sees for 7 of 10 sampled quipment (2 ultrasound equipment that uses sound nages of what is going on efibrillator/cardiac monitors - oring the rhythm of the heart; an instrument used to cut suction machines - a machine at provides suction to uid from the body) and 2) tive maintenance for 3 of 10 nedical equipment (3 nine that sterilizes medical een patient use). This failed ential to affect all patients rocedures at the clinic.	-							

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Nebraska DHHS Licensure Unit (X3) DATE SURVEY (X2) MULTIPLE CONSTRUCTION (X1) PROVIDER/SUPPLIER/CLIA STATEMENT OF DEFICIENCIES COMPLETED **IDENTIFICATION NUMBER:** AND PLAN OF CORRECTION A. BUILDING: C B. WING 08/06/2015 HC001 STREET ADDRESS, CITY, STATE, ZIP CODE NAME OF PROVIDER OR SUPPLIER 1002 WEST MISSION BELLEVUE HEALTH CENTER BELLEVUE, NE 68005 PROVIDER'S PLAN OF CORRECTION (X5) COMPLETE DATE SUMMARY STATEMENT OF DEFICIENCIES (X4) ID (EACH CORRECTIVE ACTION SHOULD BE (EACH DEFICIENCY MUST BE PRECEDED BY FULL PREFIX PREFIX CROSS-REFERENCED TO THE APPROPRIATE REGULATORY OR LSC IDENTIFYING INFORMATION) TAG TAG DEFICIENCY) G 530 G 530 Continued From page 5 Findings are: A. A tour of the clinic on 8/4/15 from 12:00 PM to G 530 7-006, 15B A-C 1:20 PM; revealed the following medical All machines have been 08/24/15 equipment with no evidence of preventive maintenance according to maintenance on the equipment: -Procedure Room 1 - ultrasound machine, Owner Manuals that have defibrillator/cardiac monitor, suction machine, and been located online or cautery machine: through the manufacturer. -Procedure Room 2 - ultrasound machine, defibrillator/cardiac monitor, and suction machine. New policies written for 08/24/15 Interview with the Director or Nursing (DON) on Machine Maintenance. 8/4/15 from 12:00 PM to 1:20 PM (during the tour) indicated that no one provided preventive The DON will check the logs maintenance on the above equipment. monthly. B. A tour of the clinic on 8/4/15 from 12:00 PM to 1:20 PM; revealed 3 autoclaves in the center See Attachments 6-23 sterilization room. The DON provided a 3-ring note book that contained log sheets titled '2015 Autoclave' for each autoclave machine. The log sheets contained an area for documenting completion of weekly, monthly and Maxi Test maintenance (a test that is completed to make sure the sterilizer is working properly). The following directions were listed at the bottom of the log sheets: "Please initial and date when completed." The log sheet for each of the 3 autoclaves only contained initials on the weekly log for January 2015. All other areas on the form were blank. C. Interview with the DON on 8/6/15 from 9:50 AM to 10:10 AM revealed that the clinic lacked a policy and procedure for preventive maintenance on equipment. Interview with the Clinic Manager 8/6/15 from 11:15 AM to 11:45 AM revealed that

the clinic had no scheduled preventive

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AND PLAN OF CORRECTION IDENTIFICATION NUMBER: A. BUILDING:	С									
HC001 B. WING	08/06/2015									
•	00/00/2015									
NAME OF PROVIDER OR SUPPLIER STREET ADDRESS, CITY, STATE, ZIP CODE										
BELLEVUE HEALTH CENTER 1002 WEST MISSION BELLEVUE, NE 68005										
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maintenance for the medical equipment.										
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