

**DEPARTMENT OF HEALTH AND HUMAN SERVICES
FOOD AND DRUG ADMINISTRATION**

DISTRICT OFFICE ADDRESS AND PHONE NUMBER 404 BNA Drive, Building 200, Suite 500 Nashville, TN 37217 (615) 366-7801 Industry Information: www.fda.gov/oc/industry	DATE(S) OF INSPECTION 03/04/2020 - 03/11/2020
	FEI NUMBER 3015826069

NAME AND TITLE OF INDIVIDUAL TO WHOM REPORT IS ISSUED TO: Kevin J. McClung, D.Ph. / Owner and PIC	
FIRM NAME Vital Care of Dickson, LLC.	STREET ADDRESS 758 Hwy 46 South, Suite 100
CITY, STATE AND ZIP CODE Dickson, TN 37055	TYPE OF ESTABLISHMENT INSPECTED Producer of Sterile Drug Products

OBSERVATION 3
 Pressure differentials are not monitored in areas where aseptic processing occurs.

Specifically,

A) Pressure differentials between areas with different air classifications (ISO 6 cleanroom and ISO 7 anteroom) are not routinely monitored/documented prior or during sterile drug production. A review of the firm's compounding records noted that the firm does not routinely document pressure differentials.

B) Pressure differentials are measured with wall-mounted manometers which are not visible from within the cleanroom. For example, I observed the firm performing sterile compounding on 03/05/2020 and I observed that the differential pressure gauge readings between the cleanroom and the anteroom were not within acceptable ranges. The cleanroom reading was .015 and the anteroom reading was .025. The firm was processing patient specific sterile drug products which included Morphine Sulfate 7.5 mg/ml and Morphine Sulfate 12 mg/ml both with a BUD of 03/05/2020 and for intrathecal administration.

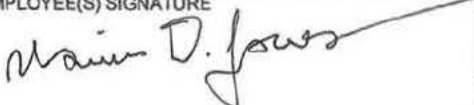
OBSERVATION 4
 Procedures designed to prevent insanitary conditions are not established or followed.

Specifically, on 03/05/2020, I observed non-sealed air gaps around the light fixtures in the firm's anteroom and cleanroom. The firm was processing patient specific sterile drug products which included Morphine Sulfate 7.5 mg/ml and Morphine Sulfate 12 mg/ml both with a BUD of 03/05/2020 and for intrathecal administration.

OBSERVATION 5
 Disinfecting agents used in the ISO 5 classified aseptic processing areas were not sterile.

Specifically, the disinfectant ((b) (4)) used by the firm is non-sterile.

Add Continuation Page

SEE REVERSE OF THIS PAGE	EMPLOYEE(S) SIGNATURE 	EMPLOYEE(S) NAME AND TITLE (Print or Type) Marvin D. Jones - Investigator	DATE ISSUED 03/11/2020
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