DEPARTMENT OF HEALTH AND HUMAN SERVICES FOOD AND DRUG ADMINISTRATION DISTRICT OFFICE ADDRESS AND PHONE NUMBER DATE(S) OF INSPECTION 06/25-29/18, 7/2, 5 & 11/18 4040 N. Central Expressway, #300 Dallas, TX 75204 FEI NUMBER 214-253-5200 3001576820 Industry Information: www.fda.gov/oc/industry NAME AND TITLE OF INDIVIDUAL TO WHOM REPORT IS ISSUED TO: David K. Johnson, Pharmacist-in-Charge FIRM NAME STREET ADDRESS CFP Acquisitions, Inc. 6136 E. 51st St. CITY, STATE AND ZIP CODE TYPE OF ESTABLISHMENT INSPECTED Tulsa, OK 74135 Producer of sterile drug products

THIS DOCUMENT LISTS OBSERVATIONS MADE BY THE FDA REPRESENTATIVE(S) DURING THE INSPECTION OF YOUR FACILITY. THEY ARE INSPECTIONAL OBSERVATIONS; AND DO NOT REPRESENT A FINAL AGENCY DETERMINATION REGARDING YOUR COMPLIANCE. IF YOU HAVE AN OBJECTION REGARDING AN OBSERVATION, OR HAVE IMPLEMENTED, OR PLAN TO IMPLEMENT CORRECTIVE ACTION IN RESPONSE TO AN OBSERVATION, YOU MAY DISCUSS THE OBJECTION OR ACTION WITH THE FDA REPRESENTATIVE(S) DURING THE INSPECTION OR SUBMIT THIS INFORMATION TO FDA AT THE ADDRESS ABOVE, IF YOU HAVE ANY QUESTIONS, PLEASE CONTACT FDA AT THE PHONE NUMBER AND ADDRESS ABOVE.

DURING AN INSPECTION OF YOUR FIRM (I) (WE) OBSERVED:

OBSERVATION 1

Personnel engaged in aseptic processing were observed with exposed skin around the face and with their face and exposed skin inside of the ISO 5 hood. Personnel did not disinfect gloves to prevent contamination.

Specifically,

- a) On 6/26/18 and 6/27/18, I observed your technician preparing sterile drug preparations in an ISO 5 laminar flow hood with exposed skin around the face (forehead, eyes and side of cheeks).
- b) On 6/26/18 and 6/27/18, I noted that your technician did not sanitize the second pair of sterile gloves after donning.

Lot #06262018@1 of Tri-Mix 15/0.5mg/5mcg/mL was made on 6/26/18. Lot #s 06272018@5 of Papaverine/ Phentolamine 30/1mg/mL, #06272018@6 of Tri-Mix 18/1/10, and #06272018@7 of Tri-Mix 30/1/10 were made on 6/27/18.

OBSERVATION 2

Media fills were not performed that closely simulate aseptic production operations incorporating, as appropriate, worst-case activities and conditions that provide a challenge to aseptic operations.

Add Continuation Page

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OF THIS
PAGE

EMPLOYEE(S) SIGNATURE

EMPLOYEE(S) NAME AND TITLE (Print or Type)

DATE ISSUED

O7/11/2018

	DEPAR	RTMENT OF HEALTH AND HUMAN SERVICES FOOD AND DRUG ADMINISTRATION		
DISTRICT OFFICE	E ADDRESS AND PHONE NUMBER	DATE(S) OF	DATE(S) OF INSPECTION	
4040 N. Cent Dallas, TX 75	ral Expressway, #300	06/25-29	06/25-29/18, 7/2, 5 & 11/18	
214-253-5200		FEI NUMBE	FEI NUMBER	
Industry Infor	mation: www.fda.gov/oc/industry	3001576	3001576820	
	OF INDIVIDUAL TO WHOM REPORT IS ISSI	UED		
TO: David K	. Johnson, Pharmacist-in-Charge			
FIRM NAME		STREET ADDRESS	PRESS	
CFP Acquisit	ions, Inc.	6136 E. 51st St.	6136 E. 51st St.	
CITY, STATE AND	ZIP CODE	TYPE OF ESTABLISHMENT INSPECTED	TYPE OF ESTABLISHMENT INSPECTED	
Tulsa, OK 74135		Producer of sterile drug products		
production, (b) (4) (b) (4)	batch sizes can be in excess of	ns or cover worst case or most challenging conditions. In routine of the performs has the operator (b) (4) This same step is then repeated (b) (4) each time.		
		'dwell time") and coverage of the item bein disinfection.	g disinfected were	
solution) or	the (b) (4) for use as a	nate contact time for the (b) (4) sporicidal. Your technician sprays the solu hood, amounting to a contact time (surface		
OBSERVA	TION 4			
Environmen	tal monitoring samples were r	not performed to ensure recovery of organis	sms.	
Specifically, (b) (4)		schnician wipe the surface of the ISO 5 lams than minutes, she performed a surface		
			Add Continuation Page	
	EMPLOYEE(S) SIGNATURE	EMPLOYEE(S) NAME AND TITLE (Print or T)	(pe) DATE ISSUED	
SEE REVERSE OF THIS PAGE	Marsaret M. C	Annes, CSO	07/11/2018	

DEPARTMENT OF HEALTH AND HUMAN SERVICES FOOD AND DRUG ADMINISTRATION DISTRICT OFFICE ADDRESS AND PHONE NUMBER DATE(S) OF INSPECTION 4040 N. Central Expressway, #300 06/25-29/18, 7/2, 5 & 11/18 Dallas, TX 75204 FEI NUMBER 214-253-5200 3001576820 Industry Information: www.fda.gov/oc/industry NAME AND TITLE OF INDIVIDUAL TO WHOM REPORT IS ISSUED TO: David K. Johnson, Pharmacist-in-Charge FIRM NAME STREET ADDRESS CFP Acquisitions, Inc. 6136 E. 51st St. CITY, STATE AND ZIP CODE TYPE OF ESTABLISHMENT INSPECTED Tulsa, OK 74135 Producer of sterile drug products OBSERVATION 5 Pressure differentials between areas with different air classifications were not monitored prior or during sterile drug production. Specifically, the magnehelic gauges that monitor the pressure differential between the ISO 7 cleanroom (room with ISO 5 laminar flow hoods) and the ISO 8 Ante Room (room where drug products are prepared) and the ISO 8 Ante Room and the unclassified pharmacy area, do not appear to be functioning properly, which may allow influx of poor quality air into a higher classified area. When the door from the unclassified pharmacy to the ISO 8 Ante Room is opened, both gauges show movement. When the door from the ISO 7 cleanroom to the ISO 8 Ante Room is opened, neither gauge moves. **OBSERVATION 6** The ISO 5 classified aseptic processing areas has difficult to clean or particle-generating equipment or surfaces. Specifically, the (b) (4) hood (ISO 5) located in your ISO 7 Cleanroom, has a cabinet that it is "mounted on a work surface constructed of pressed wood laminated with a (b) (4) Add Continuation Page EMPLOYEE(S) SIGNATURE EMPLOYEE(S) NAME AND TITLE (Print or Type) DATE ISSUED SEE REVERSE Margaret M. anna OF THIS Margaret M. Annes, CSO 07/11/2018