	F HEALTH AND HUMAN ND DRUG ADMINISTRATION			
DISTRICT ADDRESS AND PHONE NUMBER	ND DROG ADMINISTRATION	DATE(S) OF INSPECTION		
10 Waterview Blvd., 3rd Floor			2015*	
Parsippany, NJ 07054 (973) 331-4900 Fax:(973) 331-4969	rsippany, NJ 07054			
Industry Information: www.fda.gov/oc/	industry	3004600183	<u> </u>	
TO: David M. Miller, R.Ph., Owner/Pr	resident			
Millers of Wyckoff, Inc.	678 Wyckof		\$	
CITY, STATE, ZIP CODE, COUNTRY	TYPE ESTABLISHMENT IN			
Wyckoff, NJ 07481-1430 Producer of Sterile Drugs				
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 WE OBSERVED:				
OBSERVATION 1			o .	
Procedures designed to prevent microbiological contam written, and followed.	ination of drug product	s purporting to be sterile are no	t established,	
 Specifically, the following poor aseptic techniques were observed during processing on 6/29/2015 and 6/30/2015: Materials such as syringes, wrapped sterile gloves, and vials were not wiped down with Sterile (b) (4) Incomplete sanitization of the stopper of each vial of drug product with sterile (b) (4) Processing of product was performed on top of the sterile gloves wrapper, rather than the worksurface. Gloves worn during processing in the (b) (4) are not changed or sanitized with sterile (b) (4) between production of different products and/or lots. Disposable, non-sterile laboratory coats used during processing of drug products were reused, even after falling on the floor. 				
OBSERVATION 2				
Drug products do not bear an expiration date determined by appropriate stability data to assure they meet applicable standards of identity, strength, quality and purity at the time of use.				
Specifically, there is a lack of potency and sterility assurance for preserved and non-preserved sterile preparations in that beyond use dates, up to 3 months, are assigned without stability data and the container closure integrity of the bottles has not been established. Examples of products include: Buprenorphine (multidose vial) 0.3mg/mL, Lot # 02172015@24; Procaine (Buffered) (PF) 1% injectable, Lot # 04212015@17; Magnesium Sulfate Lot # 05202015@11; and Methylcobalamin(P) 25mg/mL, Lot # 04302015@20; Cefazolin Ophthalmic (PF) 50mg/mL Solution, Lot # 04212015@26; and Methylprednisone (PF) eye drops, Lot # 05152015@1.				
Additionally, there is a lack of test data to support that be (b) (4), which have been (b) (4) sterilized when produced, can (b) (4) to ensure sterility remains.				
SEE REVERSE OF THIS PAGE EMPLOYEE(S) SIGNATURE Liatte Krueger, Investig Susan M. Joseph, Investig Zakaria I. Ganiyu, Inves	TO CONTRACT	tor Liatte Knieger	07/16/2015	

	DATE(8) OF INSPECTION	
10 Waterview Blvd., 3rd Floor	06/29/2015 - 07	/16/2015*
Parsippany, NJ 07054	FEINUMBER	
(973) 331-4900 Fax: (973) 331-4969	3004600183	
Industry Information: www.fda.gov/	oc/industry	
NAME AND TITLE OF INDIVIDUAL TO WHOM REPORT ISSUED	<u> </u>	
TO: David M. Miller, R.Ph., Owner	/President	
FIRM NAME	STREET ADDRESS	
Millers of Wyckoff, Inc.	678 Wyckoff Ave	
	TYPE ESTABLISHMENT INSPECTED	
CITY, STATE, ZIP CODE, COUNTRY		

OBSERVATION 3

Each batch of drug product purporting to be sterile is not laboratory tested to determine conformance to such requirements.

Specifically,

- A. There is no documented test method for sterility testing for all products, except for Trimix (various formulations). The executed method has not been defined to be a compendial method or a method, which has been shown to be as good or better than the compendial method. In addition, on 6/29/2015, Pharmacist failed to assure us that the executed sterility testing being performed was as per method suitability conducted by a contract testing laboratory. Furthermore, negative controls are not run for each sterility test performed.
- B. Endotoxin testing is not performed on all finished lots of drug products. For example x 10mL vials, with 5mL in each vial, of Trimix 1A Lot # 04212015@10 were produced on 4/21/2015; however testing does not include endotoxin results.

OBSERVATION 4

Equipment and utensils are not maintained at appropriate intervals to prevent malfunctions and contamination that would alter the safety, identity, strength, quality or purity of the drug product.

Specifically, balance (b) (4) Model # (b) (4), which is used to weigh non-sterile ingredients has not been calibrated to ensure that the scale can accurately weigh across its minimum range (b) (4) The minimum weight used to calibrate the balance was (b) (4) in 2014 and (c) (d) in 2015. Examples of ingredients which have been weighed that are less than (c) (d) in compound procaine (b) (f) to compound procaine (b) (f) to compound procaine (b) (f) to compound procaine (b) (f) for Molybdenum (g) 25mcg/mL Lot # 06022015@5.

OBSERVATION 5

Employees are not given training in the particular operations they perform as part of their function.

Specifically,

- A. There is no documented microbiological training for the following tests and individuals who perform them:
 - Sterility and environmental testing by Pharmacist, 6 (6)
 - 2. Reading of microbiological assay results by Pharmacy Technician. (b) (6)
 - B. Performance of (b) (4) by Pharmacist Intern, (b) (6)
- B. There is no documented training for Pharmacist who performed physical testing for sterile compounded finished products.

CEE(S) SIGNATURE	DATE ISSUED
tte Krueger, Investigator, Investigator	1
an M. Joseph, Investigator	07/16/2015
	an M. Joseph, Investigator Investigator

DEPARTMENT OF HEALTH AND HUMAN SERVICES FOOD AND DRUG ADMINISTRATION				
DISTRICT ADDRESS AND PHONE NUMBER	DATE(S) OF INSPECTION			
10 Waterview Blvd., 3rd Floor	06/29/2015 - 07/16/2015*			
Parsippany, NJ 07054	FEI NUMBER			
(973) 331-4900 Fax: (973) 331-4969	3004600183			
Industry Information: www.fda.gov/oc/indu	stry			
NAME AND TITLE OF INDIVIDUAL TO WHOM REPORT ISSUED				
TO: David M. Miller, R.Ph., Owner/President	The state of the s			
FIRM NAME	STREET ADDRESS			
Millers of Wyckoff, Inc.	678 Wyckoff Ave			
CITY, STATE, ZIP CODE, COUNTRY	TYPE ESTABLISHMENT INSPECTED			
Wyckoff, NJ 07481-1430	Producer of Sterile Drugs			
C. There is no documented training for Pharmacist 60 60 who performed (b) (4) testing.				
OBSERVATION 6); X			
Aseptic processing areas are deficient regarding the system for monitoring environmental conditions.				

Specifically,

A. Procedure SOP 3.03 Environmental Monitoring (EM) of the Sterile Compounding Area (Date Effective: 01-01-07) provides instructions on surface sampling technique and incubation; however, this procedure is not being followed. For example: use of sampling environemental monitoring sampling supplies are not the same as described in the procedure, was observed to be sampling multiple locations in the (b) (4) and all EM samples are not being incubated for the (b) (4) temperature requirement.

Additionally, incubators, used for incubation of EM samples and sterility testing samples, have not been qualified and the thermometers have not been calibrated since installation of an unknown date.

- B. Non-viable particulate monitoring, performed under static conditions, is limited to (b) (4) during the certification of the (b) (4) There is no monitoring of non-viable air particulates under operating conditions within the ISO 5 (b) (4)
- C. Environmental monitoring of viable particulates is not performed within the ISO 5 (a) (4)
- D. The personnel working within the sterile area and the surfaces of the ISO 5 (b) (4) are monitored (b) (4) and not on a daily basis or after every batch of drug product produced.

OBSERVATION 7

There is a failure to thoroughly review the failure of a batch or any of its components to meet any of its specifications whether or not the batch has been already distributed.

Specifically, for at least two products which the firm has tested, including Alprostadil 10mcg/mL Lot # 01242012@1 and Methylcobalamin(P) 25mcg/mL Lot # 12042013@59, the testing for these lots resulted in a potency outside of 139% and 72%, respectively. Methylcobalamin(P) 25mcg/mL Lot # 12042013@59 was not dispensed; however the firm has produced this formulation since and has dispensed it as Lot # 06042015@37 without ensuring that the potency is within specifications.

Specifically, investigations have not been initiated for out of specification potency testing results for at least two products:

- Alprostadil 10mcg/mL Lot # 01242012@1, tested on 1/26/2012, resulted in a potency of 139%. The product was dispensed to a patient on 1/23/2012.
- Methylcobalamin(P) 25mcg/mL Lot # 12042013@59, tested on 1/16/2014, resulted in a potency of 72.2%. The lot

SEE REVERSE OF THIS PAGE	Liatte Krueger, Inve Susan M. Joseph, Inv Zakaria I. Ganiyu, I	55-00 (MD	07/16/2015
FORM FDA 483 (09/08)	PREVIOUS EDITION OBSOLETE	INSPECTIONAL OBSERVATIONS	PAGE 3 OF 5 PAGES

X		LTH AND HUMAN SERVICES UG ADMINISTRATION	Ø1
DISTRICT ADDRESS AND PHO		DATE(S) OF INSPECTION	10 Name (10)
	Blvd., 3rd Floor	06/29/2015 - 07/1 FEINUMBER	6/2015*
Parsippany, (973) 331-49		3004600183	1
Industry Info	ormation: www.fda.gov/oc/indu		
TO: David M	. Miller, R.Ph., Owner/Presid	lent street ADDRESS	100 200 200
Millers of W	Nyckoff, Inc. 678 Wyckoff Ave		
Wyckoff, NJ	07481-1430	Producer of Sterile Drugs	Ti.
	ispensed; however the firm has produced to Methylcobalamin(P) 25mcg/mL has been of	his formulation since without investigating the lispensed as Lot # 06042015@37.	e failure. Most
OBSERVATION	8	\$i	
Aseptic processing conditions.	g areas are deficient regarding the system for	or cleaning and disinfecting the equipment to	produce aseptic
Specifically, the fi	rm has not demonstrated the efficacy of the (b) (4) %, which are used to clean	the firm's (b) (4) and ISO 8 anteroom.	, Sterile (b) (4)
	id not observe personnel adhering to the es 22 Cleaning and Maintenance of the Clean	tablished contact times for their disinfectant a Room Facility (Date Effective: 01-01-07).	agents as noted in
OBSERVATION	9		10090
	e of drug product for distribution do not inc e final specifications prior to release.	clude appropriate laboratory determination of	satisfactory
Specifically,	EL.	¥	
record as per p	[생물] [[[[[[[[[[[[[[[[[[[015@31 on 6/29/2015 was not documented w Finished Preparation Testing (Date Effective:	
B. During compo check of (b) (4 control step is	(b) (4) of Phen	015@14, we observed that the operator did no colamine. There are no procedures to ensure the baserved non-dissolved particles in the (b) (4)	hat this quality
OBSERVATION	10		
	ed to prevent microbiological contamination of the sterilization process.	n of drug products purporting to be sterile do	not include
Specifically, media example, the sterile has		ne preparation of aseptically filled sterile proc 1A RX (PAPAV/PHENTOL/PGE1) 30MG/1	
hower	ver, the media fill simulation is		(b) (4)
200	EMPLOYEE(S) SIGNATURE	7/16/15 LK YK	DATE ISSUED
SEE REVERSE OF THIS PAGE	Liatte Krueger, Investigato Susan M. Joseph, Investigato Zakaria I. Ganiyu, Investiga	or Smy	07/16/2015
FORM FDA 483 (09/08)	PREVIOUS EDITION OBSOLETE INSPI	ECTIONAL OBSERVATIONS	PAGE 4 OF 5 PAGES

	DEPARTMENT OF HEAD	TH AND HUMAN S G ADMINISTRATION	ERVICES	- 11 12 - 12 - 13 - 13 - 13 - 13 - 13 -
DISTRICT ADDRESS AND PHO		G ADMINISTRATION	DATE(S) OF INSPECTION	10 N
	Blvd., 3rd Floor		06/29/2015 - 07/16/	/2015*
Parsippany,			FEI NUMBER	125
(973) 331-49		etry	3004600183	
NAME AND TITLE OF INDIVIDU	ormation: www.fda.gov/oc/indu	BLLY		
TO: David M	. Miller, R.Ph., Owner/Presid			
Millers of W	uskoff Too	STREET ADDRESS	100000	
Millers of W	yckoff, Inc.	678 Wyckoff TYPE ESTABLISHMENT INSE	PECTED	
Wyckoff, NJ	07481-1430	Producer of	Sterile Drugs	A6857 92-0-0
ODOEDWATION				250
OBSERVATION	11		197	8.
The responsibilitie	es and procedures applicable to the quality of	control unit are not	in writing and fully followed	1.
	and brockers approximate to the demand		m manag and many tono	55
	ave not qualified or audited your contract t		, used for potency, endotoxin	, and sterility
testing of Trimix f	inished products; however rely on their res	alts.		
OBSERVATION	12			8
OBSERVATION	12		W. as	
The flow of compo	onents, drug product containers, in-process	materials, and drug	g products though the buildin	ig is not designed
to prevent contami	nation.	800 80	T887 - T687 - 7	.200
Chariffeelly, the IC	10.8			0.0 (4)
which is used as th	SO 8 anteroom contains a sink sourced with (b) (4) the firm's (b) (4)	tap water that is is	ocated	(b) (4)
Willen is used us til	to (1) the limits (0) (2)	¥		
		8 D	- 1865 1865 - 1865	
				
* DATES OF INSP			ur v	
06/29/2015(Mon), 06	6/30/2015(Tue), 07/01/2015(Wed), 07/06/2015(Mon), 07/16/2015(T	'hu)	
	Na.		2	
	N			
	*		6	
	(%			
Ŧŝ				
				-
		37		
	A			
	10			
· · · · · · · · · · · · · · · · · · ·	- EMPLOYEE(S) SIGNATURE	7/16/15	K 1	DATE ISSUED
	Liatte Krueger, Investigator			
SEE REVERSE	Susan M. Joseph, Investigate		Sintle Knieger	07/16/2015
OF THIS PAGE	Zakaria I. Ganiyu, Investiga	ator	31.7	31/10/2013
FORM FDA 483 (09/08)	PREVIOUS EDITION OBSOLETE INSPE	CTIONAL OBSERV	ATIONS	PAGE 5 OF 5 PAGES