

DEPARTMENT OF HEALTH AND HUMAN SERVICES
FOOD AND DRUG ADMINISTRATION

DISTRICT OFFICE ADDRESS AND PHONE NUMBER

Division of Generic Drug Bioequivalence Evaluation
Office of Study Integrity and Surveillance USFDA-CDER
10903 New Hampshire Ave. Bldg. 51 Rm. 5308
Silver Spring, MD 20993 TEL 301-796-5032

Industry Information: www.fda.gov/oc/industry

DATE(S) OF INSPECTION

February 27 to March 3, 2017

FEI NUMBER

3007236564

NAME AND TITLE OF INDIVIDUAL TO WHOM REPORT IS ISSUED

TO: Gilbert N. Lam, President

FIRM NAME

Microconstants, Inc.

STREET ADDRESS

9050 Camino Santa Fe

CITY, STATE AND ZIP CODE

San Diego, CA 92121

TYPE OF ESTABLISHMENT INSPECTED

Bioanalytical laboratory

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:

The following observation pertains to study (b) (4) (b) (4), (b) (4) (b) (4), and (b) (4) (b) (4).

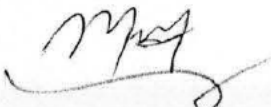
Observation 1

The neutralizing (b) (4) antibody assay cut point in studies (b) (4), and (b) (4), was not determined using samples from drug-naïve subjects.

Add Continuation Page

SEE
REVERSE
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PAGE

EMPLOYEE(S) SIGNATURE



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

Melkamu Getie-Kebtie, Ph.D., Pharmacologist
Yiyue Zhang, Ph.D., Visiting Associate

DATE ISSUED

03/03/2017

The observations of objectionable conditions and practices listed on the front of this form are reported:

1. Pursuant to Section 704(b) of the Federal Food, Drug and Cosmetic Act, or
2. To assist firms inspected in complying with the Acts and regulations enforced by the Food and Drug Administration.

Section 704(b) of the Federal Food, Drug, and Cosmetic Act (21 USC 374(b)) provides:

"Upon completion of any such inspection of a factory, warehouse, consulting laboratory, or other establishment, and prior to leaving the premises, the officer or employee making the inspection shall give to the owner, operator, or agent in charge a report in writing setting forth any conditions or practices observed by him which, in his judgement, indicate that any food, drug, device, or cosmetic in such establishment (1) consists in whole or in part of any filthy, putrid, or decomposed substance, or (2) has been prepared, packed, or held under insanitary conditions whereby it may have become contaminated with filth, or whereby it may have been rendered injurious to health. A copy of such report shall be sent promptly to the Secretary."