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 any 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:

QUALITY SYSTEM

OBSERVATION 1

The Quality Unit cannot ensure the review of all deviation reported during production prior to final product batch release.

Specifically, your Remarks Forms # GP-0004-D07 and # GP-0004-D08, ver 2, effective 12-15-2014 used to report production deviations are provided on an as needed basis by the Production Department directly to the production operators.

Traceability of these forms relies on the production operator to write in the form number and ensure all pages are included in the batch record. There are 21 cases where the production operator omitted or repeated numbering the forms. There is no assurance all Remarks Forms are included in the batch record and are reviewed by the Quality Unit prior to batch release.

In addition, the Production department determines the need for further investigation based on the deviation reported in the Remarks Form. Therefore the Quality Unit cannot ensure appropriate actions are not taken for each deviation reported in the Remark Form.
DEPARTMENT OF HEALTH AND HUMAN SERVICES
FOOD AND DRUG ADMINISTRATION

DISTRICT ADDRESS AND PHONE NUMBER:
Food and Drug Administration/CDER/OPQOS
White Oak, Building 51, Room 4316
10903 New Hampshire Avenue,
Silver Spring, MD 20993
Attn: Mr. Concepcion (Coki) Cruz
Phone: 001-301-796-3254 Fax: 001-301-847-8738

DATE OF INSPECTION: December 5-8, 2016

FIRM NAME:
Granules Omnichem Private Limited
Pharmacity SE Panwada (Mandla), Plot 121 (Part) & 122 Jawaharlal Nehru

CITY, STATE, ZIP CODE, COUNTRY:
Visakhapatnam, Andhra Pradesh, India

Establishment Inspected:
API Manufacturer

TO: Mr. Kakarlapudi Venkata Viswasatha Raja, Chief Executive Officer

OBSERVATION 2

Employee was not trained prior to conducting the cleaning of the Microbiology Laboratory ISO Class 8 room.

Specifically,

On 12/06/2016, I observed, the cleaning log book for the Microbiology Laboratory ISO Class 8 room was cleaned three times by [Redacted] who was not trained on the SOPs titled “Cleaning Procedure for Microbiology” and “Entry and Exit Procedure for Microbiology” until 5 days after first cleaning.

OBSERVATION 3

Change Control system does not evaluate and reflect all changes that may affect the production of the finished product.

Specifically,

1. Process Change Control PCR-MF1 [Redacted] did not include
   a. An adequate justification and evaluation of the addition of a [Redacted] There is no technical justification for the addition of the [Redacted] process, evaluation of the impact to the product, or specifications for the equipment to be used.
   b. Changes observed between [Redacted] [Redacted] B01-V03 and B01-V04 were not completely documented and evaluated for the following: identification of Critical Parameters, the additional, removal or change of manufacturing instructions, [Redacted] speed, and recording [Redacted] intervals.

OBSERVATION 4

SEE REVERSE OF THIS PAGE

EMPLOYEE SIGNATURE
H.L. Jamilath Selby, Investigator
Doan T. Nguyen, Investigator

DATE ISSUED: 12/9/2016
Production Master Batch Records are incomplete.

Specifically,

Your Master Batch Record for the production of [redacted] does not include the issuance of Remarks Form # GP-0004-D07 and # GP-0004-D08 ver 2, effective 12-15-2014 used to capture the following deviations, documenting additional operations, comments, observations, and addition sampling.

Some examples, but not limited to 21 omitted numbers used to identify Remarks Form for batch # [redacted] missing supervisor signatures and duplicate form numbers. Some examples of comments in these Remarks form include time period change for batch to reach required [redacted] sampling request, and change in production directions.

LABORATORY SYSTEM

OBSERVATION 5

Laboratory control records do not include complete data derived from all tests conducted to ensure compliance with established specifications and standards, including examinations and assays.

Specifically,

A p. lyrical records for the analysis of [redacted] related substances by HPLC for Batches [redacted] observed your records do not include documentation of the preparation of sample solution, stability solutions and reagents. In addition the quantity of the solution prepared, the date of preparation and the expiration date of the solution are also not documented.

FACILITIES AND EQUIPMENT
OBSERVATION 6

The facility design and maintenance increase the potential for contamination of your equipment and product.

Specifically,

On 12/5/2016, during the walk through of the production area we observed the following:

- The room used to store clean equipment contains a gap in the ceiling which spans across the length of the room that is approximately 2 inches wide exposing the material of the adjacent room and extents up to the roof.
- Cracks in the walls and peeling and chipping paint along the wall and light fixtures.
- The appearance of water damage to the drywall in various locations.

MATERIAL SYSTEM

OBSERVATION 7

Materials should be handled and stored in a manner to prevent degradation, contamination, and cross-contamination.

Specifically,

Raw material (3)(4) approved by QA on 8/10/2016 was observed being stored in the raw material warehouse which is maintained at ambient storage conditions. The labeled storage requirement for the material states “keep container cool and dry”. There is no assurance the ambient condition do not adversely affect the quality of the product. For example, during the storage period, temperatures as the log book were recorded as the following:

<table>
<thead>
<tr>
<th>Month</th>
<th>Max Temperature °C (°F)</th>
<th>Max %RH</th>
</tr>
</thead>
<tbody>
<tr>
<td>August 2016</td>
<td>34.7 (94.46)</td>
<td>89.9</td>
</tr>
<tr>
<td>September 2016</td>
<td>33.2 (91.8)</td>
<td>90.9</td>
</tr>
<tr>
<td>October 2016</td>
<td>32.2 (90.1)</td>
<td>92.0</td>
</tr>
<tr>
<td>November 2016</td>
<td>30.3 (86.54)</td>
<td>81.4</td>
</tr>
</tbody>
</table>

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See Reverse of This Page

HL Jamisah Selby, Investigator
Doan T. Nguyen, Investigator

12/9/2016