QUALITY ASSURANCE DEPARTMENT



STANDARD OPERATING	G PROCEDURE
Department: Quality Assurance	SOP No.:
Title: Core Tablet & Capsule Defect Album	Effective Date:
Supersedes: Nil	<b>Review Date:</b>
Issue Date:	Page No.:

- **1.0 Purpose:** The purpose of this SOP is to describe the action to be taken for defects observed at Core Tablets & Capsules.
- **2.0 Scope:** This SOP is applicable for the action to be taken for defects observed at Core tablet & Capsules at .....

### 3.0 References, Attachments and Annexures:

- 3.1 References:
- 3.1.1 In House
- 3.1.2 SOP: Acceptable Quality Level

### **3.2** Attachments: NA

### 3.3 Annexures:

- 3.2.1 Annexure 1: List of Defects of Core Tablet and their recommended action
- 3.2.2 Annexure 2: Defect Album for Core Tablet
- 3.2.3 Annexure 3: List of defects of Capsule & their Recommended Action
- 3.2.4 Annexure 4: Defect Album for Capsules

### 4.0 Responsibilities:

### **4.1 Quality Assurance:**

- 4.1.1 To ensure for the action to be taken for defects observed at Core tablet & Capsules.
- 4.1.1 To ensure implementation of SOP.
- 4.1.2 Concerned department Head or Designee shall be responsible for the action to be taken for defects observed at Core tablet & Capsules.

### 4.2 Regulatory Affairs, Quality Head and Plant Head:

4.2.1 To review and approve the SOP.

### **5.0 Distribution:**

- 5.1 Quality Assurance
- 5.2 Production

## 6.0 Abbreviations & Definition of terms:

### 6.1 Abbreviations:

- 6.1.1 CC No.: Change Control Number
- 6.1.2 NA. : Not Applicable
- 6.1.3 SOP : Standard Operating Procedure
- 6.1.4 MCS : Mini Capsule Sorter
- 6.1.5 DP : De-dusting and Polishing

### 6.2 Definition of terms: NA

7.0 Procedure:



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### 7.1 Action Plan to be taken for Defects Observed at Core tablet & Capsules:

- 1.1.1. Carryout the compression & capsulation as per instruction and parameters given in the BMR.
- 1.1.2. Check physical appearance of tablets & capsules during compression & capsulation process.
- 1.1.3. If defects are observed at physical appearance of the tablet & capsule during compression & capsulation, take the corrective action, by controlling the compression & capsulation process parameters and keep the minimum number of defective tablet/capsules.
- 1.1.4. Defect Album shall be maintained as per Annexure-2 & 4.
- 1.1.5. Defect should be identified during AQL Procedure.
- 1.1.6. Critical and Major defects should be considered for preparation of defect album.
- 1.1.7. Collect sample of 04 to 10 tablet/capsule from whole of the batch.
- 1.1.8. The below mentioned list of the Core Tablets/Capsule defects which are observed during Compression/Capsulation activity.

### Note:

1. If the Problem is persisting and repetitive for 03 batches then investigate the cause and to resolve the problem and review the formulation and process in consultation with F & D.



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### Annexure – 1 Tablet Defects and Recommendations

S.No.	Defects	Recommended Action
1.	Sticking	<ul> <li>Maintain temperature and RH.</li> <li>Punches and dies need polishing.</li> <li>Decrease LOD of granules.</li> </ul>
2.	Capping	<ul> <li>Increase LOD of granules.</li> <li>Remove damage punches and dies.</li> <li>Decrease machine speed.</li> </ul>
3.	Chipping	<ul> <li>Decrease the friability</li> <li>Remove damage punches or dies.</li> <li>Increase hardness.</li> <li>Handling of the tablets must be proper.</li> </ul>
4.	Weight variation	<ul> <li>Decrease machine RPM</li> <li>Granules must be uniform.</li> <li>Check granules flow parameters</li> </ul>
5.	Soft tablet	<ul> <li>Decrease machine speed</li> <li>Check granules flow property.</li> <li>Increase compressibility.</li> </ul>
6.	Oil spot	<ul> <li>Clean the punches properly.</li> </ul>
7.	Black Particle	<ul> <li>Clean the Machine.</li> <li>Check the lubricated granules.</li> </ul>
8.	Wrong Embossing	<ul> <li>Check the punches properly.</li> </ul>
9.	Picking	<ul> <li>Maintain temperature and RH.</li> <li>Decrease LOD of granule.</li> <li>Punches need polishing.</li> </ul>



### PHARMA DEVILS QUALITY ASSURANCE DEPARTMENT

STANDARD OPERATING PROCEDURE **Department:** Quality Assurance SOP No.: Title: Core Tablet & Capsule Defect Album **Effective Date:** Supersedes: Nil **Review Date:** Page No.: **Issue Date:** Annexure – 2 **Album for Tablet defects** Product:\_\_\_\_\_ Batch No.\_\_\_\_\_ **Type of Defect Type of Defect Prepared By** Checked By QA **Review By** 



QUALITY ASSURANCE DEPARTMENT

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### Annexure – 3

### Capsule Defects & Recommendation

S.No	Defects	Recommended Action
1.	Denting	<ul> <li>Ensure Proper setting of Body &amp; Cap Bush</li> <li>Ensure Proper Setting of Locking Length of capsule.</li> <li>Ensure Proper Setting of Ejection Locking Pin.</li> </ul>
2.	Telescopic/V notch	<ul> <li>Ensure Proper setting of Body &amp; Cap Bush.</li> </ul>
3.	Shade Variation/Misprint capsule	<ul> <li>Ensure the color of Empty Capsule Shells</li> <li>Ensure the printing on Empty Capsule Shell</li> </ul>
4.	Double Cap	✤ Ensure the cleaning of DP100 & MCS.
5.	Locking Length	• Ensure proper setting of machine as per BMR.
6.	Half Filled Capsule	<ul> <li>Ensure the Air Supply in Pneumatic cylinder.</li> <li>Ensure that there is no filling Hole Jamming Problem.</li> </ul>
7.	Pin Hole	<ul> <li>Ensure the Quality of Empty Capsule Shell</li> </ul>



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		Annexure -	- <b>4</b>		
	Albı	um for Capsule			
Product:			Bat	tch No	
Тур	vpe of Defect			Type of Defect	
repared By	Checked By	QA (Section Ho	ead)	<b>Review By QA Head</b>	
0 History:					
, mstory.					
			a Data	NA	
Version No.	NA	Effectiv	e Date		