

PRODUCTION DEPARTMENT

BATCH MANU				
Product Code:				
Product Name:		Generic Name:		
Document No.:	Effective	e Date:	Page No.: 1 of 19	
Batch No.:	Batch Si	ze:	Supersedes No.:	

Location:	
Block: Production Table	ets (PT)
Label Claim:	Each uncoated tablet contains: Amlodipine Besilate IP Eq. to Amlodipine
Mfg. Lic. No.:	
Product Lic. No.:	NA
Self-Life:	24 Months
MFR No.:	
Mfg. Date:	
Exp. Date:	
BMR Issued No.:	
Party:	

Issued By Stamp & Sign.

Responsibility	Name	Designation	Sign	Date
Prepared By				
Checked By				
Approved By				



PRODUCTION DEPARTMENT

### **BATCH MANUFACTURING RECORD**

Product Code: BMD		BMR No.:	
Product Name:		Generic Name: Amlodipine & Atenolol Tablets	
Document No.:	Effective Date:		Page No.: 2 of 19
Batch No:	Batch Size:		Supersedes No.: Nil

#### **1.0 MASTER FORMULA:**

#### **BILL OF RAW MATERIALS**

Sr. No.	Ingredients	Spec.	Qty. In mg Per Tablet	Overages %	Std. Qty. for 1 Lac. In Kg
Raw	V Material for Dry Mixing-				
Acti	ve Ingredients-				
1.	Atenolol	IP	50.00		5.00#
2.	Amlodipine Besilate	IP	7.25 eq. to 5.00mg		0.725#
Inac	tive Ingredients-			· · · · ·	
3.	Microcrystalline Cellulose PH-112 (MCCP-112)	IP	119.80		11.98
4.	Colour Sunset Yellow Lake	IH	1.00		0.10
5.	Sodium Starch Glycolate (Primogel)	IP	5.55		0.555
Raw	Material for Lubrication-				
8.	Sodium Lauryl Sulphate (SLS)	IP	2.00		0.20
9.	Colloidal Silicon Dioxide (Aerosil)	IP	3.00		0.30
10.	Sodium Starch Glycolate (Primogel)	IP	5.00		0.50
11.	Talcum	IP	2.00		0.20
12.	Magnesium Stearate	IP	2.00		0.20
	Weight of Uncoat	ed Tablets	200 mg		20.00 Kg

Note: # Amlodipine Besilate & Atenolol IP adds after calculation if assay below 100%.

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

#### **BATCH MANUFACTURING RECORD**

Product Code:		BMR No.:	
Product Name:		Generic Name: Amlodipine & Atenolol Tablets	
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### **CALCULATION SHEET**

#### 1- Atenolol IP is to be taken as per the formula given below:

Note: If assay of API is above 100.0% calculation not required.

### Part-A: To be calculated when single AR No.: \_\_\_\_\_\_Assay on dried basis: \_\_\_\_\_ LOD: \_\_\_\_\_

**PART-A:** To be calculated when single A.R. No. of **Atenolol IP** is to be used: If calculated quantity is less than std. qty. then dispense std. Qty.

Assay on as such basis = (100-LOD) X Assay on dried basis = \_\_\_\_\_% 100

A.R. No. of Atenolol IP	Assay on as such basis (A1)	Actual quantity of this A.R. No. to be dispensed =
	%	$\frac{\# x 100}{A1} =Kg$

PART-B: To be calculated when more than one A.R. No's of Atenolol IP is to be used:

A.R. No. of <b>Atenolol IP</b>	Assay on as such basis (A1)	Actual quantity Available (b1) (Kg)	Qty. on 100 % assay basis = (b1) x (a1) Kg 100	Remaining qty. to be dispensed $(e1) = Std. qty(c1)$
				(e1) =#
				= Kg
TOTAL (Kg)			(c1)=	

Assay of next AR. No. ----- (Assay on as such basis) (f1) = \_\_\_\_%

Actual quantity of this AR No. to be dispensed  $(g1) = \underline{(e1) \times 100} = ----Kg$ (f1)

Therefore total quantity of **Atenolol IP** to be dispensed = (b1) + (g1) =\_\_\_\_\_Kg

Assay calculation:

Sign/ Date		
Department	Done by(Production)	Checked by (Q.A.)

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

#### **BATCH MANUFACTURING RECORD**

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### **CALCULATION SHEET**

#### 2- Amlodipine Besilate IP is to be taken as per the formula given below:

Note: If assay of API is above 100.0% calculation not required.

# Part-A: To be calculated when single AR No.:\_\_\_\_\_\_Assay on dried basis: \_\_\_\_\_\_LOD: \_\_\_\_\_

**PART-A:** To be calculated when single A.R. No. of **Amlodipine Besilate IP** is to be used: If calculated quantity is less than std. qty. then dispense std. Qty.

Assay on as such basis =  $(100-LOD) \times Assay \text{ on dried basis} = ____%$ 100

100								
A.R. No. of Amlodipine Besilate IP	Assay on as such basis (A1)	Actual quantity of this A.R. No. to be dispensed =						
	%	$\frac{\# x 100}{A1} =Kg$						

**PART-B:** To be calculated when more than one A.R. No's of **Amlodipine Besilate IP** is to be used:

A.R. No. of <b>Amlodipine</b> <b>Besilate IP</b>	Assay on as such basis (A1)	Actual quantity Available (b1) (Kg)	Qty. on 100 % assay basis = (b1) x (a1) Kg 100	Remaining qty. to be dispensed (e1) = Std. qty(c1)
				(e1) =#
				(e1) =#
				= Kg
TOTAL (Kg)			(c1)=	

Assay of next AR No. ----- (Assay on as such basis) (f1) = \_\_\_\_%

Actual quantity of this AR No. to be dispensed $(g_1) = (e_1) \times 100 =Kg$
(f1)

Therefore total quantity of **Amlodipine Besilate IP** to be dispensed = (b1) + (g1) =\_\_\_\_\_Kg

Assay calculation:

Sign/ Date		
Department	Done by(Production)	Checked by (Q.A.)

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

BATCH MANU	FACTURING RECO	RD	
Product Code:	BMR	No.:	
Product Name:	Gener	Generic Name: Amlodipine & Atenolol Tablets	
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Batch No:	Batch Size:	Supersedes No.: Nil	

#### 2.0 GENERAL INSTRUCTIONS:

- Current version of SOPs should be referred during operation.
- Dispensed raw material/bulk blend/ compressed tablets should be manufactured and stored at temperature not exceeding 27°C and RH NMT 55%
- In all the processing activities, nose mask, hand gloves, secondary gown etc. shall be wearied by the personnel.
- Attach all dispensing tags and cleaning status labels with BMR.
- Clean the equipment's after use as per the standard operating procedure.
- The Blend should be compressed within 15 days.
- The compressed tablets should be packed within 30days.

#### 2.1 Line clearance of Dispensing:

Check the instructions given below and note the observation as Yes, NO or NA.

Sr. No.	Instructions	Yes/No/NA				
1.	Is dispensing area clean and free from any materials of					
2.	Whether balance is calibrated and have status label.					
3.	Scoops to be used for dispensing are clean.					
4.	LAF properly working and dispensing booth clean.					
5.	Air differential pressure, temperature and humidity with in limit (if applicable) Temp °C(NMT 27°C), RH% (NMT 55.0%), DP(6 to10 Pascal)					
6.	Material shall be least exposed to atmosphere.					
7.	Ensure proper gowning before entering to the dispensin surgical gloves shall be used while handling the materia	ıd				
Previou	Previous product name: Batch No.:					
Differential pressure across RLAF and Room:       (Limit(Between 5 to 15 Pascal))						
Checke Sign &	ed By (Production): Date:	Verified By(IPQA): Sign & Date:				

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

### BATCH MANUFACTURING RECORD

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Product Name:		Generic Name: Amlodipine & Atenolol Tablets	
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Batch No:	Batch Size:		Supersedes No.: Nil

#### **BILL OF RAW MATERIALS**

### (PRODUCTION COPY)

		Std. Qty.	@Req.	Issued		Weight in Kg		Wt. By Chkd. B			
Sr. No.	Ingredients	for 1 Lac. In Kg	Qty. in Kg	Qty. in Kg	A.R. No.	Gross	Tare	Net	Store		QA
Rav	v Material for Dry Mixing-										
Acti	ve Ingredients-						-	-		-	
1.	Atenolol IP	5.00#									
2.	Amlodipine Besilate IP	0.725#									
Inac	ctive Ingredients-			I				1	-1		
3.	Microcrystalline Cellulose PH-112 (MCCP-112) IP	11.980									
4.	Colour Sunset Yellow Lake IH	0.10									
5.	Sodium Starch Glycolate (Primogel) IP	0.555									
Rav	v Material for Lubrication-										
6.	Sodium Lauryl Sulphate (SLS) IP	0.20									
7.	Colloidal Silicon Dioxide (Aerosil) IP	0.30									
8.	Sodium Starch Glycolate (Primogel) IP	0.50									
9.	Talcum IP	0.20									
10.	Magnesium Stearate IP	0.20									

Note: # Amlodipine Besilate & Atenolol IP adds after calculation if assay below 100%.

@ Calculate the materials as per required batch size.

Dispensed by
Stores
Date

Checked by Production Date Verified by QA Date Page No. 7 of 19 store copy

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

### BATCH MANUFACTURING RECORD

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Product Name:		Generic Name: Amlodipine & Atenolol Tablets		
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Batch No:	Batch Size:		Supersedes No.: Nil	

### **BILL OF RAW MATERIALS**

### (STORE COPY)

		Std. Qty.	@Req.	Issued		W	eight in	Kg	Wt. By	Chkd. By	
Sr. No.	Ingredients	for 1 Lac. In Kg	Qty. in Kg	Qty. in Kg	A.R. No.	Gross	Tare	Net	Store		QA
Rav	v Material for Dry Mixing-										
Acti	ve Ingredients-						-				
1.	Atenolol IP	5.00#									
2.	Amlodipine Besilate IP	0.725#									
Inac	ctive Ingredients-										
3.	Microcrystalline Cellulose PH-112 (MCCP-112) IP	11.980									
4.	Colour Sunset Yellow Lake IH	0.10									
5.	Sodium Starch Glycolate (Primogel) IP	0.555									
Raw	v Material for Lubrication-										
6.	Sodium Lauryl Sulphate (SLS) IP	0.20									
7.	Colloidal Silicon Dioxide (Aerosil) IP	0.30									
8.	Sodium Starch Glycolate (Primogel) IP	0.50									
9.	Talcum IP	0.20									
10.	Magnesium Stearate IP	0.20									

Note: # Amlodipine Besilate & Atenolol IP adds after calculation if assay below 100%.

@ Calculate the materials as per required batch size.

Dispensed by	
Stores	
Date	

Checked by Production Date Verified by QA Date

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

### **BATCH MANUFACTURING RECORD**

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### 2.2 Weight Verification Sheet:

Balance ID: \_\_\_\_\_

Sr.		~		Std.		Issued Qty.	Checked By	Verified	
No.	Ingredients	Spec.	UOM	Quantity (Kg)	A.R. No.	Gr. wt.	(Production)	by (IPQA)	
MAT	FERIAL FOR GRANULATI	ON:							
1.	Atenolol	IP							
2.	Amlodipine Besilate	IP							
3.	Microcrystalline Cellulose PH-112 (MCCP-112)	IP							
4.	Colour Sunset Yellow Lake	IH							
5.	Sodium Starch Glycolate	IP							
MAT	FERIAL FOR LUBRICATIO	DN:							
1.	Sodium Lauryl Sulphate	IP							
2.	Colloidal Silicon Dioxide	IP							
3.	Sodium Starch Glycolate	IP							
4.	Talcum	IP							
5.	Magnesium Stearate	IP							

### 3.0 GRANULATION PROCESS:

Granulation started on:\_\_\_\_\_

#### 3.1 Line clearance of Granulation:

Previous product: \_\_\_\_\_

Batch No.:\_\_\_\_\_

	Cleaning done by:, Cleaning done by:,	eaned On:		,
Sr. No.	Instructions	Yes/No/NA	Checked By (Production)	Verified By (IPQA)
1	Ensure that all equipment and utensils are clean and dry and status board affixes (Record as per Table-1).			
2	Is area free from any materials of previous batch?			
3	Whether the container, sieve, scoops and auxiliary items are cleaned.			
4	Check the room temperature. TempºC (NMT 27°C) and Differential pressure Pascal (6 to 10 Pascal).			
5	AHU system under operation or not.			
6	Calibration status of Equipment/instrument complies or not.			
7	Balance calibration status is OK or not.			
8	Whether swab/rinse sample testing report complies or not? (if applicable)			
9	Whether the wall, floor and light in satisfactory condition?			

.

	Prepared By	Checked By	Approved By
Signature			
Date			



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### **BATCH MANUFACTURING RECORD**

Product Code:	BI	AR No.:		
Product Name:	Ge	Generic Name: Amlodipine & Atenolol Tablets		
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Batch No:	Batch Size:	Supersedes No.: Nil		

### EQUIPMENT STATUS CHECKLIST

Sr. No.	Name of Equipment	Equipment ID No.	Observation (Should be clean and dried)	Checked (Production)	Verified By (IPQA)
1.	Sifter		Yes/No		
2.	Octagonal blender		Yes/No		
3.	Balance		Yes/No		
4.	SS scoop		Yes/No		

**3.2** Sifting: Sift separately the following material and collect in poly bags/containers. Check sieve integrity before and after use.

#### SIFTING OF GRANULATION MATERIALS

Ingredient	Qty. In	Sieve	Sieve In	From	То	Done By/	Ckd. By/	
ingreulent	Kg	Size (#)	Before Use	After use	FIOII	10	Date	Date
Atenolol IP								
Amlodipine Besilate IP								
Microcrystalline Cellulose PH-112 IP								
Colour Sunset Yellow Lake IH								
Sodium Starch Glycolate IP								

### SIFTING OF BLENDING MATERIAL / LUBRICANTS

		Sieve	Sieve In	tegrity		То	Done	Ckd By/ Date
	Qty. In Kg	Size (#)	Before Use	After use	From		By/ Date	
Sodium Lauryl Sulphate IP								
Colloidal Silicon Dioxide IP								
Sodium Starch Glycolate IP								
Talcum IP								
Magnesium Stearate IP								

	Prepared By	Checked By	Approved By
Signature			
Date			



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### 3.3 MANUFACTURING PROCESS:

Step No.	Manufacturing Instruction	Eq. ID.	From	То	Done By/ Date	Ckd. By/ Date
3.3.1	Dry Mixing:					
	Atenolol (#Kg), Amlodipine Besilate (#Kg),         Microcrystalline Cellulose PH-112 (Kg), Sunset         Yellow Lake (Kg) and Sodium Starch Glycolate         (Kg) in blender and mix for 20 minutes.					
3.3.2	Blending & Lubrication:					
	Add Sodium Lauryl Sulphate (Kg), Colloidal Silicon Dioxide (Kg), Sodium Starch Glycolate (Kg) and Talcum (Kg) in blender with dry mixing granules and mix it for 20 minutes.					
	Add <b>Magnesium Stearate</b> (Kg) in blender and mix for further for 5 minutes.					

### 3.4 GRANULE WEIGHING RECORD:

Container No.	Gross wt. (Kg)	Tare wt. (Kg)	Net wt. (Kg)	Done By/ Date	Ckd By/ Date
1/					
2/					
3/					
4/					
5/					
6/					
7/					
8/					
9/					
10/					
Total					

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

BATCH MANU	FACTURING	RECORD		
Product Code:		BMR No.:		
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#### 3.5 SAMPLING OF BLEND:

• After completion of the manufacturing activity batch card shall be checked by production executive and inform IPQA through analytical request after completion of granulation process.

**Checked By (Production)** 

• IPQA shall review batch card and visually inspect of the material for physical Appearance, labeling status, number of container etc. and will collect the sample as per SOP, and shall submit to QC for analysis.

Checked By (IPQA)

• After release from QC, IPQA shall paste the 'APPROVED" label on each container.

#### **3.6 YIELD RECONCILIATION:**

A = Theoretical batch size	= Kg / tablets
B = Actual quantity of blend	= Kg
C = Samples	=

D = Yield = B / A x 100

(Note: - Granulation yield NLT 99.00%)

Loss Quantity:

Checked by (Production): Date: Verified by (QA): Date:

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

### BATCH MANUFACTURING RECORD

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Batch No:	Batch Size:		Supersedes No.: Nil	

#### 4.0 COMPRESSION:

Date: \_\_\_\_\_

Started At:	

### 4.1 Line clearance:

### Previous product: \_\_\_\_\_\_, Batch No.: \_\_\_\_\_\_

Sr. No.	Instructions	Observations	Checked (Production)	Verified By (IPQA)
1	Is area free from any materials of previous batch?	Yes/No		
2	Whether area and utensils cleaned?	Yes/No		
3	Whether the compression machine is cleaned and set as per SOP and have "CLEANED" label affixed? Equipment ID No.:	Yes/No		
4	Check the room temperature, RH and differential pressure =°C (NMT 27°C), RH= % (NMT 55%). Differential PressurePascal (6 to 10 Pascal)	OK/NOT OK		
5	All the equipment shall be used during process are cleaned.	Yes/No		

#### 4.2 Process:

Sr. No.	Compression Instruction	Observations	Checked (Production)	Verified By (IPQA)
1.	Collect the approved granules from the granules store for compression.			
2.	Ensure the correct punch set is assembled in the compression machine.			
3.	Ensure the availability and online filling of Batch Document.			
4.	Collect the tablets as per total no. of punches from each side and check them individually for any damages on upper and Lower Surface before continuing the operation of compression machine. Check and Record the observation and details of die & punch in the table A: <b>Die and punch verification</b>			
5.	If compression time is less than one hour, minimum Three observations shall be recorded.			
6.	Ensure that all the data of actual processing are entered in log book of individual equipment/Instrument.			
7.	Collect the compressed tablets in polythene lined container. Weight the containers and record the weights in table given below, label them properly and transfer them to bulk store (Container number should be given as $1/x$ , $2/x$ where x is the total number of containers			

	Prepared By	Checked By	Approved By
Signature			
Date			



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### BATCH MANUFACTURING RECORD

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Batch No:	Batch Size:		Supersedes No.: Nil		

#### Table: A-Die and punch verification

	Punch Specification																							
	Туре				B-To	Γooling & Stations																		
Punch Details         Upper Punches         8.00 mm (Flat Round Break line )           Lower Punches         8.00 mm (Flat Round plain )											Б		0 10 -											
											– Dies : 8.10 mm													
	Punch	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23
Upper	No.																							
Punches																								
	Punch	24	25	26	27	28	29	30	31	32	33	34	35											
	No.																							
	Punch	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	19	20	21	22	23
	No.																							
Lower Punches																								
runches	Punch	24	25	26	27	28	29	30	31	32	33	34	35											
	No.																							

Checked by (Production):\_\_\_\_\_

Verified By (IPQA):\_\_\_\_\_

#### 4.3 IN PROCESS CHECKS:

#### 4.3.1 Specification:

Sr. No.	Parameters	Requirement	Frequency of Monitoring
1.0	Description	Flat Round break line one side plain on other side	At the start of machine
2.0	Weight of 20 tablets	4.00gm <u>+</u> 3%	Every 30 Minutes
3.0	Avg. weight	200 mg <u>+</u> 3%	Every 2 Hours
4.0	Uniformity of weight	200 mg <u>+</u> 5%	Every 2 Hours
5.0	Thickness $4.40 \pm 0.2 \text{ mm}$ Every 2 Hours		Every 2 Hours
6.0	Hardness	NLT 2.0 Kg/cm <sup>2</sup>	Every 2 Hours
7.0	Friability	NMT 1%	Every 2 Hours
8.0	DT	NMT 15 min	Every 2 Hours
9.0	Diameter	8.10 mm <u>+</u> 0.2 mm	At the start of machine
10.0	Appearance	Orange colour round shape tablets.	Every 2 Hours
11.0	Temperature	NMT 27℃	Every 2 Hours
12.0	RH	NMT 55%	Every 2 Hours

	Prepared By	Checked By	Approved By
Signature			
Date			



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	BATCH	I MANU	FACTU	RING R	ECORD						
Product Code:				H	BMR No.	:					
Product Name:				(	Generic N	Name: Ai	nlodipine	e & Ateno	olol Tabl	ets	
<b>Document No.:</b>			Effecti	ive Date:		Р	age No.:	14 of 19			
Batch No:			Batch	ch Size: Supersedes No.: Nil							
4.4 In-process observation	on sheet fo	r producti	ion:								
Description:											
Diameter:											
	Date										
Wt. of 20 Tabs.	Time										
4.00gm <u>+</u> 3%	LHS										
4.00gm <u>+</u> 378	RHS										
	Date										
Wt. of 20 Tabs.	Time										
4.00gm <u>+</u> 3%	LHS										
<u>.</u>	RHS										
	Date										
Thickness	Time										
4.40 <u>+</u> 0.2 mm	LHS										
	RHS										
	Date										
Friability	Time										
(NMT 1.0 %)	LHS										
	RHS										
Hardness	LHS										
(NLT 2.0 Kg/cm <sup>2</sup> )	RHS										
DT	LHS										
NMT 15 min.	RHS										
Appearance: Orange colour flat	LHS										
round break line one other side.	RHS										
Temperature (NMT 27°C)											
RH (NMT 55%)											
Done By											

Attached additional sheet if required...

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

		BATCH MAN	UFACTUR	ING RECO	RD						
Product (	Code:			BMR	No.:						
Product N	Name:		-	Gener	ic Name:	Amlodipine &	Atenolol Ta	ablets			
Document	t No.:		Effectiv	e Date:		<b>Page No.:</b> 15	of 19				
Batch No:	:		Batch S	ize:	Supersedes No.: Nil						
			WEIGHT V.	ARIATION C							
Average W	Average Weight of Tablet:     Frequency     Even										
Date:											
Time:											
1.											
2.											
3.											
4.											
5.											
6.											
7.											
8.											
9.											
10.											
11.											
12.											
13.											
14.											
15.											
16.											
17.											
18.											
19.											
20.											
Avg. Wt.											
Min wt.											
Max wt.											
Checked by											

Attached additional sheet if required......

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

BAT	CH MANUFACTURING RECORD	
Product Code:	BMR No.	:
Product Name:	Generic N	Name: Amlodipine & Atenolol Tablets
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Batch No:	Batch Size:	Supersedes No.: Nil

### 4.5 In-process observation sheet for IPQA:

Description:						
Diameter :						
	Date					
Wt. of 20 Tabs.	Time					
4.00gm <u>+</u> 3%	LHS					
	RHS					
	Date					
Wt. of 20 Tabs.	Time					
4.00gm <u>+</u> 3%	LHS					
	RHS					
	Date					
Thickness	Time					
4.40 <u>+</u> 0.2 mm	LHS					
	RHS					
	Date					
Friability	Time					
(NMT 1.0 %)	LHS					
	RHS					
Hardness	LHS					
(NLT 2.0 Kg/cm <sup>2</sup> )	RHS					
DT	LHS					
NMT 15 min.	RHS					
<b>Appearance:</b> Orange colour flat	LHS					
round break line one other side.	RHS					
Temperature (NMT 27°C)						
RH (NMT 55%)						
Done By						

Attached additional sheet if required......

	Prepared By	Checked By	Approved By
Signature			
Date			



	PRODUCTION DEPARTMENT								
		BATCH MAN	UFACTUR	ING RE	CORD	)			
Product	Code:			B	MR No	.:			
Product	Name:			Ge	eneric I	Name: Amlo	dipine & Ater	nolol Tablets	
Documer	nt No.:		Effective	e Date:		Page	e No.: 17 of 19	)	
Batch No	:		Batch Si	ze:		Supe	ersedes No.: N	Nil	
WEIGHT VARIATION OF 20 TABLETS									
Average V	Veight of Table	et:			Fr	requency	ŀ	Every 2 hours.	
Date:									
Time:									
1.									
2.									
3.									
4.									
5.									
6.									
7.									
8.									
9.									
10.									
11.									
12.									
13.									
14.									
15.									
16.									
17.									
18.									
19.									
20.									
Avg. Wt.									
Min wt.									
Max wt.									
Checked by									

Attached additional sheet if required......

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

BATCH MANU	FACTURING	RECORD		
Product Code:				
Product Name:		Generic Name: Amlodipine & Atenolol Tablets		
Document No.: Effective Da		ate: Page No.: 18 of 19		
Batch No:	Supersedes No.: Nil			

#### 4.6 TABLET WEIGHING RECORD:

Container No.	Gr. wt.	Tare wt.	Net wt.	Container No.	Gr. wt.	Tare wt.	Net wt.
1/				11/			
2/				12/			
3/				13/			
4/				14/			
5/				15/			
6/				16/			
7/				17/			
8/				18/			
9/				19/			
10/				20/			
Total net weight	of Tablets:						
Checked By(Sig	n & Date):						

### 4.7 SAMPLING :

After completion of the manufacturing activity batch card shall be checked by production executive and inform IPQA through analytical request after completion of compression process.

### Checked By (Production)

- IPQA shall review batch card and then visually inspect the bulk for physical appearance, labeling status, number of container etc. and will collect the sample as per SOP, and shall submit to QC for analysis.
- After release from QC IPQA shall paste the **'APPROVED**" label on each drum.

#### 4.8 VISUAL INSPECTION OF TABLET:

Date: \_\_\_\_\_

Time Duration		Quantity rejected	Done by	
From	То	Quantity rejected	Done by	

Total weight of rejected tablets: \_\_\_\_\_ Good Tablet weight: \_\_\_\_\_

% Yield: \_\_\_\_\_

Checked by (Production): (Sign & Date)

#### Verified by (IPQA): (Sign & Date)

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

BATCH MANU	FACTURING	RECORD		
Product Code:				
Product Name: Generic Name: Amlodipine & Atenolo				l Tablets
Document No.:	<b>Page No.:</b> 19 of 19			
Batch No: Batch Size:			Supersedes No.: Nil	

### **4.9 YIELD RECONCILIATION:**

•	Average weight of tablets (A)=: mg	
٠	Total weight of uncoated tablets (B) = Kg.	
•	В	
	Quantity of uncoated tablet in Number (C)=	X 1000 X1000 =
	A	
•	Samples (D)=	
•	C + D	
	Yield= 100 =	(NLT 98.00%)
	Actual batch size	
Chec	ked By (Production):	Verified By (IPQA):

### 5.0 FINAL REVIEW OF BATCH CARD ON SHOP FLOOR:

Production manager/Designee shall review the batch card will give his comment, if any.

Checked By (Prod. Mgr.)

#### **5.0 ANY DEVIATION:**

 Deviation No.
 Reason for deviation

Checked By (Prod. Manager)

#### 7.0 HISTORY SHEET:

BMR No.	New BMR No.	Revision No.	Reason of revision
		00	New BMR

	Prepared By	Checked By	Approved By
Signature			
Date			