

PRODUCTION DEPARTMENT

### BATCH PACKING RECORD

<b>Product Code:</b>		BPR No.:			
Product Name:			Generic Name: Atenolol Tablets IP 25 mg		
<b>Effective Date:</b>				<b>Page No.:</b> 1 of 24	
Batch No.:		Batch	Size:	Supersedes No.: Nil	
Location:					
Block: Production Tablets	(PT)				
Label Claim:	Atenolol IP	ted tablet cor 25 n q.s.	ng		
Mfg. Lic. No.:					
Product Lic. No.:	NA				
Self-Life:	mon	ıths			
Pack Style:	20x14 Table	ets			
Country Name:	Domestic				
Change Control No.:	NA				
Mfg. Date:					
Exp. Date:					
BMR ISSUED NO.:			_		
MRP:					
		Issued	l By Stamp & Sign.		

Responsibility	Name	Designation	Sign	Date
Prepared By				
Checked By				
Approved By				



PRODUCTION DEPARTMENT

#### BATCH PACKING RECORD

<b>Product Code:</b>	BPR No.:		
Product Name:	Generic Name: Atenolol Tablets IP 25 mg		
<b>Effective Date:</b>		<b>Page No.:</b> 2 of 24	
Batch No.:	Batch Size:	Supersedes No.: Nil	

#### 1.0 GENERAL INSTRUCTIONS:

- ➤ Good manufacturing practices should be followed during the entire process of packing.
- All the Equipments used for packing should be properly cleaned as per the relevant SOP.
- All the Equipments and containers should have proper status label with Stage, Product name, B. No., Mfg. Date etc.
- All the equipments should be operated as per the relevant SOP's only.
- Issued packing materials should be cross checked by production personnel against dispensing sheet before taking up for packing.
- ➤ Overwriting in BPR shall be strictly avoided & correcting shall be made as per SOP.
- > All the activities should be carried out according to the BPR only. All the operations shall be carried out in clean and orderly manner.
- Any deviation in process shall be bought to knowledge of QA and prior approval of QA department should be taken.
- > Critical parameters like temperature, Humidity and pressure differences should be checked and monitored.
- ➤ In process controls should be carried out throughout the packing operations as per relevant BPR and relevant SOP's.
- > Ensure that all the packing materials, in process materials and finished goods should be placed in respective areas with proper label to avoid mix up.
- Attach additional issue sheets from QA, wherever required.
- Attach system generated data sheets wherever applicable.

	Prepared By	Checked By	Approved By
Signature			
Date			



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Product Code: BPR No.:				
Product Name:	Generic Name: Atenolol	Generic Name: Atenolol Tablets IP 25 mg		
<b>Effective Date:</b>		<b>Page No.:</b> 3 of 24		
Batch No.:	Batch Size:	Supersedes No.: Nil		

#### 2.0 DISPENSING OF PACKING MATERIALS:

#### 2.1Instructions:

- 1. Follow the packing materials dispensing SOP.
- 2. Appropriate weighing balances should be used while issue.
- 3. Ensure that weighing balances are calibrated & Verified on daily basis.
- 4. Printed Al. Foil and Special /PVC should be issued in poly bags.
- 5. Each roll should be labeled separately.
- 6. Cartons should be issued in bundles.
- 7. Cartons should be kept in plastic/shippers crates covered with lid or supplier and properly labeled.
- 8. Carton should be closed with transparent Cello tape.
- 9. One complaint slip is pasted on inside flap of corrugated box.
- 10. Shippers should be issued in bundles with proper label.
- 11. Keep all issued materials on separate pallets in PM dispensing room.

#### 2.2 Line Clearance Checks:

Line clearance Given By(IPQA):

Sign & Date

Sr. No.	Line Clearance Checks	Observation	Checked by QA
1.	Containers used for previous batch/product removed from area		
2.	All status labels of previous batch/products are removed		
3.	BPR or any other documents related to the previous batch / product are removed from area.		
4.	Absence of any previous product /batch materials.		
5.	Cleanliness of the area		
6.	Cleanliness of the area below balances/ pallets.		

#### 2.3 Line clearance certificate for area and equipment:

Area	PM dispensing room	Equipment	Weighing Balance
Area Cleaned By:		Equipment No.:	
Checked By:		Equipment Cleaned By:	
Previous Product:		Batch No.:	
Checked By (Packing Su	pervisor):		
Sign & Date			

	Prepared By	Checked By	Approved By
Signature			
Date			



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<b>Product Code:</b>	BPR No.:		
Product Name:	Generic Name: Atenolol	Generic Name: Atenolol Tablets IP 25 mg	
<b>Effective Date:</b>		<b>Page No.:</b> 4 of 24	
Batch No.:	Batch Size:	Supersedes No.: Nil	

#### 2.4 BILL OF PACKING MATERIALS:

(BPR Copy) Dispensed on: \_\_\_\_\_

Sr.	Items	Qty. for 1 Lac.	*Req. Qty. In	Issued Qty. In	A.R. No.	Issued by	Chec By	
No.	Teens,	In Kg/Nos.	Kg/Nos.	Kg/Nos.	11.11.11.11.	Store	Prod.	QA
1	<b>Printed Foil</b> Foil Width = 198mm	2.50 Kg						
2	<b>Base Foil:</b> Clear Transparent PVC film, Foil Width:204mm	12.00 Kg						
3	<b>CARTON</b> –Dim: 100x 38x96 mm (20X14 Tablets)	358 Nos.						
4	<b>5 PLY CORRUGATED BOX-</b> Dim (OD): 425(L)x405(W)x 391(H) mm, (160 Cartons per box 4x5x6) Mkt.by address is printed in corr. box length panel in red colour.	3 Nos.						
5	<b>BOPP TAPE -</b> BOPP Pre Printed 48 mm x 65 mtrs.	1 Nos.						
6	Cello tape ½ inch	1 Nos.						

Note: \*Calculate the materials as per required batch size.

Dispensed By: Checked By: Verified By: (Store) (Prod. Supervisor) (QA)

Store copy page No.: 5 of 22

	Prepared By	Checked By	Approved By
Signature			
Date			



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<b>Product Code:</b>		BPR No.:				
Product Name:		Generic Name: Atenolol Tablets IP 25 mg				
<b>Effective Date:</b>			Page No.	: 5 of 24		
Batch No.:	Batch S	Size:	Supersec	les No.: Nil		

#### **BILL OF PACKING MATERIALS**

(STORE COPY) Dispensed on: \_\_\_\_\_

Sr.	Items	1 Lac.	rkeq. Qıy. In	Qıy. m	A.R. No.	Issued by	Checked By	
No.	TOTAL D	In Kg/Nos.	Kg/Nos.	Kg/Nos.	111111111111	Store	Prod.	QA
1	<b>Printed Foil</b> Foil Width = 198mm	2.50 Kg						
2	<b>Base Foil :</b> Clear Transparent PVC film, Foil Width:204mm	12.00 Kg						
3	<b>CARTON</b> –Dim: 100x 38x96 mm (20X14 Tablets)	358 Nos.						
4	<b>5 PLY CORRUGATED BOX-</b> Dim (OD): 425(L)x405(W)x 391(H) mm, (160 Cartons per box 4x5x6) Mkt.by address is printed in corr. box length panel in red colour.	3 Nos.						
5	<b>BOPP TAPE -</b> BOPP Pre Printed 48 mm x 65 mtrs.	1 Nos.						
6	Cello tape ½ inch	1 Nos.						

Note:\*Calculate the materials as per required batch size.

Dispensed By: Checked By: Verified By: (Store) (Prod. Supervisor) (QA)

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

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<b>Product Code:</b>		BPR No.:				
Product Name:		Generic Name: Atenolol Tablets IP 25 mg				
<b>Effective Date:</b>			<b>Page No.:</b> 6 of 24			
Batch No.:	Batch S	Size:	Supersedes No.: Nil			

#### 3.0 PACKING SPECIFICATION:

Sr. No.	Description	Over Printing Matter Standards (For Example only)	Over Printing Matter Actual	Chec By						
<b>A.</b>	Primary Packi	• • • • • • • • • • • • • • • • • • • •		Prod.	QA					
1.	Blister			1	т——					
	Blister coding details	B. No MFG EXP M.R.P. Rs PER 14 TABS.INCL OF ALL TAXES								
В.	Secondary Pac	king:		-11						
	Unit Carton	t Carton Printed								
	Carton details	20 x14 Tablets								
1.	Carton coding details	Batch No.: Mfg. Date: Exp. Date: MRP Rs.: (Incl. of all Taxes) Per 14 Tablets								
C.	Tertiary Packi	ng								
	5 ply shipper	5 ply printed shipper								
	Shipper details	160 cartons in one 5 ply shipper								
1.	Shipper coding details	B.No. MFG. EXP. Qty. 160 X 20 X 14 TABS.								
	Sealing of Shipper/BOPP Tape	Printed BOPP Tape in "H" type on top and bo	ttom.							

### 3.1 STANDARD PACKING INSTRUCTIONS:

- Check and verify the status board/label.
- All the materials of previous batches should be removed and line clearance certificate to be obtain from IPQA before starting any activity.
- Transfer the QC Released Tablets of the Batch to the primary cubicle.

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

#### BATCH PACKING RECORD

Product Code:	BPR No.:	BPR No.:				
<b>Product Name:</b>	Generic Name: Atenolol T	Generic Name: Atenolol Tablets IP 25 mg				
<b>Effective Date:</b>		<b>Page No.:</b> 7 of 24				
Batch No.:	Batch Size:	Supersedes No.: Nil				

- Produce the blister of 2x14 tablets using 198 mm printed aluminum foil & 204 mm base foil on a blister packing machine. The blister foil should be duly overprinted with the respective batch legend.
- Blister sealing leak test should be performed periodically to monitor the sealing.
- Each blister should be visually inspected to reject the defective ones.
- 20x14 such inspected blister should be packed inside each printed carton. The carton should be duly overprinted with the respective batch legend.
- 160 such inspected unit cartons should be packed inside the each shipper.
- The shipper should be properly labeled using coder. The coding details should be overprint with the respective batch legend on the shipper label.
- Each shipper should be sealed using Pre-printed BOPP tape in "H" type on top and bottom.
- After completion of the batch packing, intimate IPQA department through the transfer ticket.
- Complete the BPR for reconciliation of the batch after that transfer the packed shippers to the Finish Goods Store.

3.2	PACKING -	Date:
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#### **Instructions:**

- a. Gowning should be follows as per SOP.
- b. Masks and gloves should be used in the primary packing.
- c. Check for the cleanliness of the area and equipment.
- d. Check the Temperature, Humidity, and differential Pressure as per BPR or as per SOP
- e. Check that batch/product is released by QC for packing before starting of packaging operations and transfer to primary packing.
- f. Check the status label on the area on the display board outside the packing cubical.
- g. Operate blister packing machine as per SOP.
- h. Line clearance should be given take during any shift change.
- i. Line clearance procedure should also be followed in case of change in stereo or any major breakdown which can affect the packing quality.

	Prepared By	Checked By	Approved By
Signature			
Date			



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	BATCI	H PACKING	G REC	ORD							
Product Code: BPR No.:											
Product Name:				Generic	Name	: Atenolo	ol Tab	lets IP	25 mg		
	ive Date:								<b>o.:</b> 8 of 2		
Batch	No.:	В	atch S	ize:			Sı	uperse	des No.	: Nil	
	ne clearance check (Initial/shearance of Packing Line	O	,	ase Tick √	If Yes &	Σ X If No	or Not	Applic	cable	1	T
Sr.	Clearance Checks	Date									
No.		Time									
1.	Product name:										
2.	Area Cleanliness below/ Bal etc.	ance/ Pallets/									
3.	Machine Cleanliness										
4.	Packaging material of previous remove.	ous product									
5.	Over coding details on bliste	er									
Sr.		Date									
No.	Clearance Checks	Time									
6.	Over coding details on unit	carton									
7.	Pasting cello tape										
8.	Over coding details on outer	carton									
9.	Product Packaging Insert										
10.	O. Specimen of 5 Ply Shipper coding										
11.	11. Correctness of status label										
12. Daily Verification of balances											
Check	Checked by Production (Sign/Date)										
Verifi	ed by IPQA (Sign/Date)										
3.4 Ve	rification of Tablet received	from core ar	ea:	<u> </u>	L	1		1	1	1	

Total Container No.	Total Weight	Checked by Production	Verified by IPQA

### 3.5 Stereo detail:

Issue the required number of stereos to operator and retrieve the same from them after completion of activity and record shall be maintained as per table given below;

	stereos from QA		reos given erator	No. of stereos returned by Operator		Total No. of stereos submitted to QA		Submitted by	Retrieved
Carton	Blister	Carton	Blister	Carton	Blister	Carton	Blister	(Packing)	By (IPQA)

	Prepared By	Checked By	Approved By
Signature			
Date			



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<b>Product Code:</b>		BPR No.:	
<b>Product Name:</b>		Generic Name: Atenolol T	Tablets IP 25 mg
<b>Effective Date:</b>			<b>Page No.:</b> 9 of 24
Batch No.:	Batch S	Size:	Supersedes No.: Nil

#### 3.6 Line clearance overprinting of carton:

- Line clearance of the area and machine.
- ii. Affix the specific batch stereo and prepare a specimen proof for the approval of packing supervisor and then by IPQA supervisor & affix in the BPR.
- iii. After approval start coding of carton and check the each carton for correctness and legibility of the batch detail. iv. In-process, rejection and destruction of rejected cartons shall be recorded.

Line clearance certificate for area and equipment					
Area:	Equipment:	Carton coding machine			
Area Cleaned By:	Equipment No.:				
Checked By:	Equipment Cleaned By:				
Previous Product:	Batch No.:				
Checked By (Packing Supervisor): Sign & Date					
Line clearance Given By(IPQA): Sign & Date					

#### Over coding detail for Blister, carton and shipper

	Over printi				
Sr. No.	Details on PM (for example)	Actual details	Blister	Carton	Shipper
1					
2	Batch No.:				
3	Mfg. Date:				
4	Exp. Date:				
5	M.R.P.: (Incl. of all taxes) Per 14 Tablets				
6	Qty. 160 X 20 X 14 TABS.				
Do alviu a	Signature				
Packing	Packing Date				
TDO A	Signature				
IPQA	Date				

**Note:** Which is not applicable mention NA and put tick mark which is applicable.

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

#### BATCH PACKING RECORD

<b>Product Code:</b>	BPR No.:	
Product Name:	Generic Name: Atenolo	Tablets IP 25 mg
<b>Effective Date:</b>		<b>Page No.:</b> 10 of 24
Batch No.:	Batch Size:	Supersedes No.: Nil

#### 3.7 Reconciliation of Packing Material:

Sr. No.	Particulars	Cartons	Shipper
1	Quantity Issued		
2	Quantity coded		
3	Good inspected quantity		
4	Quantity rejected		
5	Qty. destroyed		
6	Qty. destroyed by		
7	Checked by Prod. (Sign/Date)		
8	Verified by IPQA (Sign / Date)		

#### 3.8 Shipper coding:

- i. Arrange the klass marker of respective batch no. for coding on unit carton and arrange the alphabets for shipper label coding as per information given in the BMR and first take a specimen on carton and shipper label coding specimen on plain A4 size paper & get the approval from packing supervisor and then from IPQA.
- ii. After approval all the unit carton/shipper of the batch shall be coded and if any unit carton/shipper rejected during coding same shall be destructed and record shall be maintained.

#### 4.0 BLISTER PACKING:

#### 4.1 Machine Setting:

1. Take line clearance from IPQA.

Line clearance certificate for area and equipments:								
Area	Equipment	Blister Machine						
Area Cleaned By	Equipment No.							
Checked By	Equipment Cleaned By							
Previous Product	Batch No.							
Checked By (Packing Supervisor): Sign & Date								
Line clearance Given By(IPQA): Sign & Date								

- 2. Check the change parts as per product specification.
- 3. Mount the rollers and check the cavity alignment of sealing roller.
- 4. Mount BCP, and affix stereos.
- 5. Adjust forming & sealing temperature and pressure.
- 6. Load the printed and plain foil, and adjust machine to smooth foil run and take out proof of Batch coding. Get the approval from packing supervisor and IPQA.
- 7. Set the sealing temperature 180°C to 200°C. Forming Temp150°C to 160°C.
- 8. Ensure proper Knurling and cutting length.
- 9. Check status label on Tablets containers.

	Prepared By	Checked By	Approved By
Signature			
Date			



#### PRODUCTION DEPARTMENT

#### BATCH PACKING RECORD

<b>Product Code:</b>	BPR No.:			
Product Name:	Generic Name: Atenolol 7	Generic Name: Atenolol Tablets IP 25 mg		
<b>Effective Date:</b>		<b>Page No.:</b> 11 of 24		
Batch No.:	Batch Size:	Supersedes No.: Nil		

- 10. Load the hopper with Tablets to be stripped.
- 11. Operate the blister packing machine as per SOP.
- 12. Check the leak test of blister as per Leak Test SOP. Record it in in-process control record.
- 13. Attach approved specimen sample to BPR duly signed by Packing Supervisor and QA Personnel.

#### **4.2** General instruction:

- 1. Carry out blistering operation after batch printing approval by production supervisor & IPQA.
- 2. Record the parameters at a stated frequency.
- 3. Carry out the Leak test as per SOP.
- 4. Note the changes in foil rolls and splices.
- 5. Check the coding on each splice and foil at the start and end. Check at least 1 meter section of each side.
- 6. Foil rolls / Splices should be numbered.
- 7. Attach the sample of every new foil roll and every splice in each roll with BPR.
- 8. Note the Machine start, stop and end time.

#### 4.3 Blister Packing Start up Control Checks:

- 1. Run the machine and collect few initial blisters.
- 2. Check for Knurling, Cutting, sealing, batch overprinting, etc. and observation shall be recorded.
- 3. If the initial parameters are satisfactory, continue packing.
- 4. In process test observation shall be recorded both by packing and IPQA supervisor as per table No.4.4
- 5. Reasons for machine stop should be recorded. In the following tables.

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Signature			
Date			



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<b>Product Code:</b>		BPR No.:				
Product Name:	Generic Name: Atenolol T	Tablets IP 25 mg				
<b>Effective Date:</b>			<b>Page No.:</b> 12 of 24			
Batch No.: Batch S		ize:	Supersedes No.: Nil			

#### 4.4 Secondary and tertiary packing:

- 1. Pack the number of blister in carton then followed by outer carton and finally in shipper as per requirement given in section 2.0 (packing specification).
- 2. Each carton and shipper shall weigh to identify the shortage if any.
- 3. Close the shipper by BOPP tape properly.
- 4. Person involve in the packing shall be recorded as per following table:

Date				
Time	To	То	То	То
Inspection of blister done by				
Counting of blister done by				
Carton				
Insertion of blister & Carton done by				
Inspection of over coding on carton done by				
Shipper coding done by				
Insertion of Carton in shipper done by				
Shipper sealed and weighed by				
Checked by:				
Production/packing				
IPQA				

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

### **BATCH PACKING RECORD**

<b>Product Code:</b>		BPR No.:			
Product Name:		Generic Name: Atenolol Tablets IP 25 mg			
<b>Effective Date:</b>			<b>Page No.:</b> 13 of 24		
Batch No.:	Batch S	Size:	Supersedes No.: Nil		

### 5.0 IN PROCESS CHECK:

### 5.1 In-process check by production at initial and every 30 min.

Sr.	In process	Date						
No.	checks	Time						
1.	Temp.							
2.	RH							
3.	Forming rolle temperature							
4.	Sealing roller Temperature							
5.	Check working of NFD by removing one tablet from each track							
6.	Tab. with fore black particle	eign /						
7.	Foil shifting							
8.	Batch detail o	n foil						
9.	No. of tab/ Bl	ister						
10.	Proper cutting blister	g of						
11.	Leak test (Hourly)							
12.	Proper gluing carton							
13.	No. of blister printed carton	1						
14.	Batch detail of printed carton	1						
15.	Seal the carto cello tape							
16.	No. of carton shipper							
17.	Batch details on shipper label							
18.	Pasting of BC	PP tape						
Check	xed by (Produc	ction)						

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

### **BATCH PACKING RECORD**

<b>Product Code:</b>		BPR No.:					
Product Name:		Generic Name: Atenolol Tablets IP 25 mg					
<b>Effective Date:</b>			<b>Page No.:</b> 14 of 24				
Batch No.: Batch S		Size:	Supersedes No.: Nil				

### In-process check by production at initial and every 30 min.

			n-process	check by	production	)11 at 1111	tiai aiiu	cvery.	ж ппп.		
Sr. No.	In process checks	Date Time									
1.	Temp.	Time									
2.	RH										
2.											
3.	Forming rolle temperature										
4.	Sealing roller Temperature										
5.	Check workin NFD by remo one tablet from track	ving									
6.	Tab. with fore black particle										
7.	Foil shifting										
8.	Batch detail o	n foil									
9.	No. of tab/ Bl	ister									
10.	Proper cutting blister	g of									
11.	Leak test (Hourly)										
12.	Proper gluing carton										
13.	No. of blister printed carton										
14.	Batch detail o printed carton										
15.	Seal the cartor cello tape										
16.	No. of carton shipper	in one									
17.	Batch details shipper label	on									
18.											
Check	xed by (Produc	ction)									

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

### **BATCH PACKING RECORD**

<b>Product Code:</b>		BPR No.:					
Product Name:		Generic Name: Atenolol Tablets IP 25 mg					
<b>Effective Date:</b>			<b>Page No.:</b> 15 of 24				
Batch No.:	Batch S	Size:	Supersedes No.: Nil				

### In-process check by production at initial and every 30 min.

			ii-pi ocess	 ~J P-	ouucu	 	 		
Sr. No.	In process checks	Date Time							
		11me							
1.	Temp.								
2.	RH								
3.	Forming roller temperature								
4.	Sealing roller Temperature								
5.	Check workin NFD by remo one tablet from track	ving							
6.	Tab. with fore black particle	ign /							
7.	Foil shifting								
8.	Batch detail o	n foil							
9.	No. of tab/ Bl	ister							
10.	Proper cutting blister	gof							
11.	Leak test (Hourly)								
12.	Proper gluing carton								
13.	No. of blister printed carton								
14.	Batch detail o printed carton								
15.	Seal the cartor cello tape								
16.	No. of carton shipper								
17.	Batch details on shipper label								
18.	Pasting of BOPP tape								
Check	xed by (Produc	ction)							

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

### **BATCH PACKING RECORD**

<b>Product Code:</b>		BPR No.:					
Product Name:		Generic Name: Atenolol Tablets IP 25 mg					
<b>Effective Date:</b>			<b>Page No.:</b> 16 of 24				
Batch No.:	Batch S	Size:	Supersedes No.: Nil				

### In-process check by production at initial and every 30 min.

	I		-process cne	CK Dy p	Toducuo	, at 1111	uai aiiu	every.	, IIIII.		
Sr.	In process	Date									
No.	checks	Time									
1.	Temp.										
2.	RH										
3.	Forming rolle temperature										
4.	Sealing roller Temperature										
5.	Check working NFD by remoone tablet from track	ving									
6.	Tab. with fore black particle										
7.	Foil shifting										
8.	Batch detail o	on foil									
9.	No. of tab/ Bl										
10.	Proper cutting blister	g of									
11.	Leak test (Hourly)										
12.	Proper gluing carton	of									
13.	No. of blister printed carton										
14.	Batch detail of printed carton										
15.	Seal the carto cello tape										
16.	No. of carton shipper										
17.	Batch details shipper label	on									
18.											
Check	ked by (Produc	ction)									

Attach additional sheet if required....

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

### BATCH PACKING RECORD

<b>Product Code:</b>		BPR No.:				
Product Name:		Generic Name: Atenolol Tablets IP 25 mg				
<b>Effective Date:</b>			<b>Page No.:</b> 17 of 24			
Batch No.:	Batch S	Size:	Supersedes No.: Nil			

### 5.2 In-process check by IPQA for initial and every 60 min

Sr.	In process	Date						
No.	checks	Time						
1.	Temp.							
2.	RH							
3.	Forming roller temperature	r						
4.	Sealing roller Temperature							
5.	Check workin NFD by remo- one tablet from track	ving n each						
6.	Tab. with fore black particle	ign /						
7.	Foil shifting							
8.	Batch detail on foil							
9.	No. of tab/ Bl							
10.	Proper cutting blister	of						
11.	Leak test (Bi-hourly)							
12.	Proper gluing carton							
13.	No. of blister printed carton							
14.	Batch detail of printed carton							
15.	Seal the cartor cello tape							
16.	No. of carton shipper							
17.	Batch details of shipper label	on						
18.	Pasting of BOPP tape							
Check	ked by (IPQA)							

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

### BATCH PACKING RECORD

<b>Product Code:</b>		BPR No.:	
Product Name:		Generic Name: Atenolol T	Tablets IP 25 mg
<b>Effective Date:</b>			<b>Page No.:</b> 18 of 24
Batch No.:	Batch S	Size:	Supersedes No.: Nil

### In-process check by IPQA for initial and every 60 min

Sr.	In process	Date						
No.	checks	Time						
1.	Temp.							
2.	RH							
3.	Forming rolle temperature							
4.	Sealing roller Temperature							
5.	Check workin NFD by remo one tablet from track	ving						
6.	Tab. with fore black particle	eign /						
7.	Foil shifting							
8.	Batch detail o	n foil						
9.	No. of tab/ Bl							
10.	Proper cutting blister	g of						
11.	Leak test (Bi-hourly)							
12.	Proper gluing carton							
13.	No. of blister printed carton	L						
14.	Batch detail o printed carton							
15.	Seal the cartor cello tape							
16.	No. of carton shipper							
17.	Batch details shipper label							
18.	Pasting of BC	PP tape						
Check	ked by (IPQA)							

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

### **BATCH PACKING RECORD**

<b>Product Code:</b>		BPR No.:	
Product Name:		Generic Name: Atenolol T	ablets IP 25 mg
<b>Effective Date:</b>			<b>Page No.:</b> 19 of 24
Batch No.:	Batch S	Size:	Supersedes No.: Nil

### In-process check by IPQA for initial and every 60 min

	process encours, in Quarter minus over, ou min										
Sr.	In process	Date									
No.	checks	Time									
1.	Temp.										
2.	RH										
3.	Forming roller temperature	r									
4.	Sealing roller Temperature										
5.	Check workin NFD by remo one tablet from track	ving									
6.	Tab. with fore black particle	ign/									
7.	Foil shifting										
8.	Batch detail o	n foil									
9.	No. of tab/ Bl										
10.	Proper cutting blister	of									
11.	Leak test (Bi-hourly)										
12.	Proper gluing carton	of									
13.	No. of blister printed carton										
14.	Batch detail o printed carton										
15.	Seal the cartor cello tape										
16.	No. of carton shipper										
17.	Batch details of shipper label	on									
18.	Pasting of BO	PP tape									
Check	ked by (IPQA)										

Attach additional sheet if required....

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

	BATCH	PACKING REC	CORD						
<b>Product Code</b>	:		BPR No.:						
<b>Product Name</b>	2:		Generic Name: Atenolol Tablets IP 25 mg						
<b>Effective Date</b>	<b>:</b>		-1	Page No.: 20 of 24					
Batch No.:		Batch S	Size:		Superse	des No.: Nil			
	VEIGHING RECORD		17						
	filled shipper:					T			
Shipper No.	Gross wt. In Kg.	Weighing done		No. Gross	wt. In Kg.	Weighing done by			
1.			20.						
2.			21.						
3.			22.						
4.			23.						
5.			24.						
6.			25.						
7.			26.						
8.			27.						
9.			28.						
10.			29.						
11.			30.						
12.			31.						
13.			32.						
14.			33.						
15.			34.						
16.			J						
17.									
18.									
19.									
	FY7 • 1 4		3.6 G1:	***					
Min. Shipper			Max. Shij	pper Weight:	, p. (TDC)				
Chec	cked By (Production S	supervisor)		Veril	y By (IPQ	<b>A</b> )			
Loose Shipper N	No.:								
••	<del></del>								
	Prepare	d By	Checked	l By	A	pproved By			
Signature	•			•		•			
Date									



PRODUCTION DEPARTMENT

$\mathbf{B}^{A}$	ΥT	$\mathbf{CH}$	PA	CKING	RECORD	)
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<b>Product Code:</b>	BPR No.:			
<b>Product Name:</b>	Generic Name: Atenolol Tablets IP 25 mg			
<b>Effective Date:</b>		<b>Page No.:</b> 21 of 24		
Batch No.:	Batch Size:	Supersedes No.: Nil		

#### 7.0 RECONCILIATION OF PACKING MATERIAL:

Sr. No.	Material	Printed Aluminum foil	Base foil	Printed Cartons	<b>BOPP Tape</b>	Shippers
1.	Std. Qty.					
2.	Quantity Issued					
3.	Extra Qty. issued					
4.	Qty. used					
5.	Qty. returned (attach MRN)					
6.	Qty. destroyed after coding					
7.	Qty destroyed after pkg.					
8.	Total qty. destroyed					
9.	Qty. destroyed by					
10.	Checked by Prod. (Sign/date)					
11.	Verified by IPQA (Sign/date)					
12.	Remarks					

### 8.0 FINISH PRODUCT SAMPLING AND QUALITY CONTROL APPROVAL:

Production person shall raise the sample request and provide to IPQA for sampling. IPQA shall perform sampling as per respective SOP and sent to QC.

Requisition raised By (Packing Supervisor): _	Sampled By (IPQA): _	
Sampling Details:		

S.No.	Sample detail	Date	Quantity	Sampled By			
1.	Sample for analysis						
2.	Control Samples						
3.	Stability Samples						
4.	Other Sample						
	Total Oty, of samples=						

	Prepared By	Checked By	Approved By
Signature			
Date			



	PRODUCTION DEPARTMENT								
		BATCH PACKI	NG REC	ORD					
Produ	Product Code: BPR No.:								
Produ	uct Name:			Generic N	Name: Atenolol T	Cablets IP	25 mg		
Effec	tive Date:					Page No	.: 22 of 24		
Batch	n No.:		Batch S	ize:		Superse	des No.: Nil		
Tr <b>D</b> a	ansfer finished g	OS TRANSFER TO FG goods to FG Stores. Throu			each a copy of T.T.	to BPR			
	No. of shippers p	acked							
	er shipper								
	blister per Carto								
~ •	f Tablets transfer								
- •	f shippers transfe	erred to BSR							
Transf	er note No.								
Sign o	f Packing Super	rvisor							
Sign o	f BSR Supervis	or							
	ATCH RECON	CILIATION:							
Sr. No.		Particulars	5		In K	Σg	In No.		
1.									
2.	2. Partial								
3.	3. Packing loss (Non recoverable)								
4.	4. Quantity actually transferred to FG Store								
5.	Sample:								
5a.	Analysis Samp	le Qty.							
5b.	Control Sample	es Qty.							
5c.	Stability Sample Qty.								

### **Reconciliation of Batch Yield:**

Total packed Quantity (4+5a+5b+5c+5d)

Other Sample Qty.

Accountability=

5d.

6. 7.

Yield	=	Total Quantity Packed (6) + Partial x 100 Batch size
	=	x 100
	=	% (NLT 97.0 %)

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

### BATCH PACKING RECORD

<b>D</b> A	ATCH PACKII	NG RECORD				
<b>Product Code:</b>		BPR No.:				
Product Name:		Generic Name: A	Generic Name: Atenolol Tablets IP 25 mg			
<b>Effective Date:</b>	1			<b>(o.:</b> 23 of 24		
Batch No.:		Batch Size:	Supers	edes No.: Nil		
Remark:						
(Packing Superviser)			(IPQA)			
11.0 DEVIATION APPROV	'AL:					
Deviation No.	Reason for devi	iation				
12.0 REVIEW OF BPR:			Date:			
Particulars		Status		Checked By QA		
Signature of Authorized Person	ns					
<b>Contents and Enclosures:</b>		1	1			
PM Requisition						
PM Issue Order						
Excess material issue note, if a	ny					
PM return note (if applicable)						
Specimens of Packing material						
In Process packing control rep	orts					
TR of Finished Product Pack						
COA of Finished Product						
FG Goods Transfer Note						
Final Dispatch Note						
Destruction and approvals						
Deviation and its Justification						
Reconciliation and Yields						
Legibility of contents						

	Prepared By	Checked By	Approved By
Signature			
Date			



PRODUCTION DEPARTMENT

		BATCH PACE	KING RECOR	D	
Prod	uct Code:		BF	PR No.:	1
Produ	uct Name:		Ge	neric Name: Atenolol	Γablets IP 25 mg
Effec	tive Date:				<b>Page No.:</b> 24 of 24
Batch	n No.:		<b>Batch Size:</b>		Supersedes No.: Nil
13.0	DISPATCH AD	VICE:			
	D 1 4		FOR THE USE		T
	Product: Batch No:				
	Qty. Released: A.R			A.R. No:	:
	Released Date:				
	The BPR has be	en reviewed and the	above batch is r	eleased for Dispatch.	
	Signature of QA	Manager/Designee	:	Date:	
14.	HISTORY SHE	ET:			
	DDD M.	N DDD N	D M .	D	Character IN

BPR No.	New BPR No.	Revision No.	Reason of revision	Change Control No.
		00	-	

	Prepared By	Checked By	Approved By
Signature			
Date			