

National Quality Control Laboratory

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SAMPLE INFORMATION FORM

Date Sample Submitted:	Labora	atory Reference N	Io:	
Product Generic/Brand Na	me:			
Product Chemical Name:				
Product Descrip	tion:			
Product Presenta	tion:			
Label c	laim: ————			
Batch/Lot No: Date of manufacture:		Product License Date of Ex	e No:	
Name of Client and				
Country of Origin:	Samp	les ed:		nples rned
Test(s) requested: a) Identification b) Dissolution c) Assay d) Uniformity of Weight e) f)	Limits:	U.S.P B.P Ph. Eur		and exact page):
Analyst: Mary Magda	Signature:		Date:	19-02-2015
Checked by:	Signature:		Date:	
Approved by:	Signature:		Date:	

UNIFORMITY OF WEIGHT: TABLETS/CAPSULES/SACHETS/VIALS

No.	Tablets/Capsules/ Sachets/Vials (mg)	Empty Capsule/ Sachet/Vial (mg)	Capsule/Sachet/Vial Content (mg)	% Deviation From mean (for deviating tabs/caps)
1				• • •
2				
3				
4				
5				
6				
7				
8				
9				
10				
11				
12				
13				
14				
15				
16				
17				
18				
19				
20				
Total:				
Avg:				
Calcula Deviation				

Comments:			

ASSAY DATA FORM

Standard Preparation for Assay:	ASSAY
Sample Preparation for Assay:	

DISSOLUTION

	Standard Pre	paration	for D	issolution
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Dissolution Conditions			
	1st Run	2 nd Run	3 rd Run
Dissolution Medium:			
Volume used:			
Apparatus:			
Rotations per minute:			
Time (min)			

Describe below any subsequent dilutions after the dissolution:

CHROMATOGRAPHIC CONDITIONS:

	ASSAY		
Column No:	Type of Column:		
Column Temp (°C):			
Detection λ (nm):	Injection Vol (μL):		
Mobile Phase: Composition (% v/v)	& Ratios	Flow Rate (mL/min):Pump Pressure (bars):	
Column No:	DISSOLUTION Type of Column:		
Column Temp (°C):			
Detection λ (nm):	_ Injection Vol (μL):		
Mobile Phase: Composition (% v/v) &	& Ratios	Flow Rate (mL/min):Pump Pressure (bars):	
REFERENCE SUBSTANCES:			

NO	Reference Substances/Related Substances	NQCL Code/Batch	Purity (%)
1.			
2.			
3.			
4.			
5.			

	REAGENTS USED						
			Lot/Batch	Date	Expiry		
	Reagent Name	Manufacturer	No.	Opened	Date	Remarks	
1.							
2.							
3.							
4.							
5.							
6.							
7.							
8.							

	EQUIPMENT USED						
	Equipment Name	NQCL No./Code	Date of Last Calibration	Date of Next Calibration	Remarks		
1.							
2.							
3.							
4.							
5.							
6.							
7.							
8.							

APPENDIX

Describe in Summary the reagent preparation procedures including mobile phase and l

Report any other tests carried out on the sample.

	WORKSHEET TRACKING							
No.	ACTIVITY	FROM: OFFICER/ ANALYST	SIGNATURE	TO: OFFICER/ ANALYST	SIGNATURE	DATE		
1								
2								
3								
4								
5								
6								
7								