

National Quality Control Laboratory

Hospital Road , KNH Complex, P.O. Box 29726, 00202 Nairobi, Kenya Telephone: 2726963, +254 - 020 - 3544525/30 • Fax: 2718073 Email: info@nqcl.go.ke Website: www.nqcl.go.ke

SAMPLE INFORMATION FORM

Date Sam	nple Submitted: _	2015-01	Labora	tory Reference	No: MULT	TICOMP
Product (Generic/Brand Na	me: Pred	nisolone Tablets			
Product (Chemical Name:	<u>Pred</u>	nisolone			
	Product Descrip	otion:				
	Product Presenta	ation:				
	Label o	rlaim: Pred	imisolone			
	Batch/Lot No: _ of manufacture: _ ne of Client and _	122365 1970-01-01 Chemonics Ke	nya Pharma LLC	Product Licer Date of l		01-01
Cl:	Address: _	P.o Box 1325 (00606 Nairobi, Kenya			
Cher	nt Reference No: _ Manufacturer:	MYLAN LABS	LTD			
Со	untry of Origin: _	INDIA	Sample	es d: 50 Capsules	Sam Retu	nples rned
Test(a) b) c) d) e) f)	(s) requested: Identification Dissolution Assay Uniformity of Weight pH		Limits:	Monograph U.S.P _ B.P Ph. Eur Ph. Intl Other's _		and exact page):
Analyst:	Mary Magda	ı	Signature:		Date:	19-02-2015
Checked	by:		Signature:		Date:	
Approve	d 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: Column Temp (°C):	DISSOLUTION Type of Column:		
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									