## ""INDUSTRIAL TRAINING""

# A REPORT SUBMITTED TO SAVITRIBAI PHULE PUNE UNIVERSITY, PUNE



FOR THE DEGREE OF MASTER OF SCIENCE

IN

ORGANIC CHEMISTRY

UNDER THE FACULTY OF SCIENCE

BY

Mr. Saurabh Laxman Sanap

Department of Chemistry, G. M. D.Arts, B.W. Commerce and

Science College, Sinnar

UNDER THE GUIDANCE OF

Prof. :- Dr. M.R. Gaware

Head of

## DEPARTMENT OF CHEMISTRY

G.M.D.ARTS, B.W.COMMERCE AND SCIENCE COLLEGE,

SINNAR 422103

APRIL 2023







Maratha Vidya Prasarak Samaj's G.M.D. ARTS, COMMERCE AND SCIENCE COLLEGE, SINNAR, DISTRICT- NASHIK DEPARTMENT OF CHEMISTRY (PG)

## CERTIFICATE

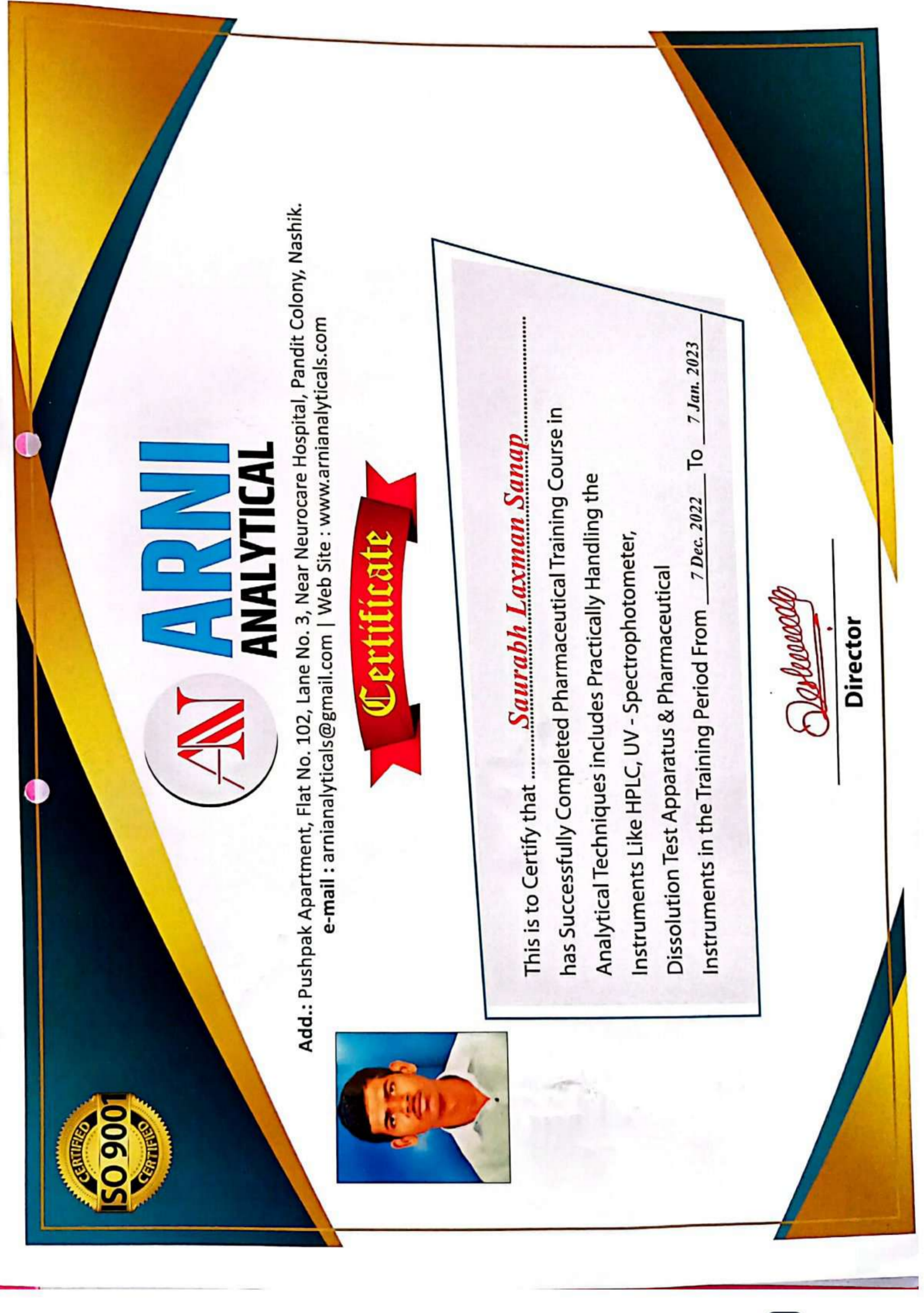
This is to certify that Mr. Saurabh Laxman Sanap studying in M.Sc.-II (Organic Chemistry) at M.V.P. Samaj's G.M.D. Arts, B.W. Commerce and Science College, Sinnar has successfully completed "Pharmaceutical Training Course in Analytical Techniques" (CHO-453-Industrial Training) from 07/12/2022 to 07/01/2023 conducted by Arni Analyticals, Nashik during the semester IV of academic year 2022-2023.

HOD Chemistry

DEPARTMENT OF CHEMISTRY G.M.D. Arts, B.W. Commerce and Science college. Sinnar

Principal

G.M.D.Arts, B.W.Commerce and Science College, Sinner, Dist. Neshik







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## Certificate

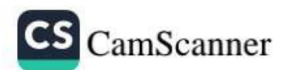
"Pharmaceutical Training Course in Analytical Techniques"

This is to certify that Mr./Miss/ Mrs. Saurabh Laxman Sanap studying in M. Sc.-II (Organic Chemistry) at M. V. P. Samaj's G. M. D. Arts, B. W. Commerce and Science College, Sinnar has successfully completed "Pharmaceutical Training Course in Analytical Techniques" from 07/12/2022 to 07/01/2023 conducted by Arni Analyticals, Nashik and has obtained "B+" grade.

Mr. Masum Deshmukh

Director

Add.: Pushpak Apartment, Flat No. 102, Lane No. 3, Near Neurocare Hospital, Pandit Colony, Nashik. | e-mail: arnianalyticals@gmail.com





## **ACKNOWLEDGEMENT**

The success and final outcome of this training required a lot of guidance and assistance from many people. All that I have done is only due to such supervision and assistance and I would never forget to thank them.

I respect and thank Respected Dr. P.V. Rasal Sir for providing me an opportunity to do the training and giving all the support and guidance which made me complete the training successfully. I am extremely thankful to him for providing such a nice support and guidance.

I owe my deep gratitude to Prof. Manoj Gaware Sir (Head of Chemistry Department) who took interest on my training and guided me all along, till the completion of training by providing all the necessary information.

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Name: - Saurabh Laxman Sanap





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## TENELIGLIPTIN

### Introduction-

- Teneligliptin is a pharmaceutical drug for the treatment of type-2 diabetes mellidus.
- Teneligliptin belongs to the category of medicines called "anti-diabetic".
- It is used along or in combination with other drugs to lower blood sugar levels.
- Teneligilptin tablet contains the teneligilptin which belongs to class of dipeptidyl peptidase-4 inhibitors.
- It works by blocking the action of DPP-4 (an enzyme that destroys the harmone 'Incretin'). The
  enzyme 'Incretins' helps to produce more insulin only when required and reduces the liver's blood
  sugar level when not needed.

### Chemical Formula- C22H30N6O5

## Molar Mass- 426.58 gm/mol

- Teneligliptin significantly controls glycemic parameters with safety. No dose adjustment is required.
- As we all know that teneligiptin tablet contains only 20 mg active ingredient i.e. teneligiptin.
   Other layers or coatings are exceplents.
- Once a tablet is formulated then directly it doesn't comes to market. First of all some of the random tablets are collected and forwarded for testing.

## Testing have 2 types-

- 1. Physical
- 2. Chemical

## **Physical Testing-**

- 1. Average weight test
- 2. Uniformity of weight
- 3. Thickness
- 4. Dimensions
- 5. Hardness

## **Chemical Tests-**

- Dissolution Test
- 2. Separation Technique (HPLC)
- 3. Absorbance

## Structure of Teneligliptin-





## FINISHED PRODUCT SPECIFICATION AND TEST METHOD

NAME OF PRODUCT: TENELIGLIPTIN TABLETS 20 MG

PAGE NO .:- Page 2 of 5

#### TEST METHOD

1) Description: White coloured, round shaped, film coated tablets, plain on both sides.

Identification:

The retention time of the principal peak in the chromatogram of sample preparation should correspond to that of the standard preparation as obtained in the "Assay".

3) Average weight:

Weigh together 10 tablets selected at random and calculate the average weight.

Calculation:

Average weight (mg) = -----10

Where, W= Weight of 10 tablets in mg

Limit:  $283 \text{ mg} \pm 7.5\%$ 

4) Uniformity of Weight:

Select randomly 10 tablets and weigh individual tablet. Calculate average, the minimum and maximum value.

Limit: 283 mg ±7.5% (Between 261.8 mg and 304.2 mg)





## FINISHED PRODUCT SPECIFICATION AND TEST METHOD

NAME OF PRODUCT: TENELIGLIPTIN TABLETS 20 MG

PAGE NO .:- Page 3 of 5

### 5) Dissolution (By HPLC):

Dissolution Pa	aram	eters :				
Medium	:	Water	Rotatory Speed	:	75 rpm	
Volume	1:	900 mL	Temperature	:	37°C ± 0.5°C	
Apparatus	:	USP Type II (Paddle)	Time	:	45 Minutes	

### 11.1 Preparation of Solutions:

### Standard preparation:

Weigh and transfer accurately about 22 mg of Teneligliptin (Equivalent to 32.43 mg Teneligliptin Hydrobromide Hydrate) working standard to a 100 mL volumetric flask add 70 mL of water and sonicate to dissolve and make up the volume with water.

#### Sample preparation:

Pour 900 mL of dissolution medium in each vessel. Allow sufficient time for the dissolution medium to equilibrate at  $37^{\circ}\text{C} \pm 0.5^{\circ}\text{C}$ . Adjust stirring element speed to 75 rpm. Place one tablet in each of the six vessels and immerse the paddles in the dissolution medium so that there is a distance of  $25 \text{mm} \pm 2 \text{mm}$  between the bottom of the paddle and inside bottom of the vessel. Start the apparatus.

At the end of specified time intervals (after 45 minutes), withdraw 10 mL aliquot from a zone midway between the surface of the dissolution medium and the top of the rotating paddle and filter through 0.45µ filter paper discarding first few mL of the filtrate. Inject directly.

#### 11.2 Procedure:

Measure the absorbance of the resulting solution at 210nm.

#### Calculations:

Teneligliptin

(% Drug Release) = 
$$\frac{At}{As} \times \frac{Ws}{100} \times \frac{900}{LC} \times \frac{P}{100} \times \frac{426.57}{628.86} \times 100$$

Where,

A<sub>T</sub> = Absorbance due to Teneligliptin in the sample preparation.

As = Absorbance due to Teneligliptin in the standard preparation.

Ws = Weight of Teneligliptin hydrobromide hydatre working standard taken for standard preparation, in mg.

P = Purity of Teneligliptin hydrobromide hydatre working standard used for standard

LC = Label claim of a tablet, in mg. 426.57 = Molecular weight of Teneligliptin

628.86 = Molecular weight of Teneligliptin hydrobromide hydatre

Limits: Not less than 80.00 % of labeled amount is dissolved in 45 minutes





## FINISHED PRODUCT SPECIFICATION AND TEST METHOD

NAME OF PRODUCT: TENELIGLIPTIN TABLETS 20 MG

PAGE NO .:- Page 4 of 5

### 13) Assay (By HPLC):

#### Reagents Required:

Sr.No.	Name of Reagent	Grade
1	Water	HPLC grade
2	Acetonitrile	HPLC grade
3	Octane-1-sulphonic acid sodium salt	AR grade
4	O-Phosphoric acid	AR grade

#### 13.1 Chromatographic Conditions:

Column	:	C18, (150 mm X 4.6 mm), 5µm
Pump mode	:	Isocratic
Mobile Phase	:	Buffer: Acetonitrile (60:40)
Flow rate	:	1.0 mL/min
Injection volume	:	20 μl
Column Temperature	:	30°C
Wavelength	:	UV, 210 nm
Run time	:	1.5 times of the retention time of principle peak

## 13.2 Preparation of Mobile Phase:

## Preparation of Buffer:

Dissolved 0.1M Potassium dihydrogen orthophosphate in 1000 mL of water; Prepare a mixture of Buffer, Acetonitrile (60:40 v/v), filter through 0.45 µ filter and degas.

#### 3.3 Preparation of solutions:

## Standard preparation:

Weigh and transfer accurately about 20 mg of Teneligliptin (29.48 mg Teneligliptin Hydrobromide Hydrate) working standard to a 100 mL volumetric flask add 70 mL of water and sonicate to dissolve and make up the volume with water.

#### Sample preparation:

Weigh 10 tablets and determine average weight. Crush the tablets to a fine powder. Weigh and transfer powder equivalent to 20 mg of Teneligliptin to a 100 mL dry volumetric flask. Add 70 mL of water, sonicate for not less than 20 minutes with intermittent shaking. Make up the volume with water. Filter through 0.45  $\mu$  Nylon filter discarding first few mL of the filtrate.

#### 13.4 Evaluation of System Suitability:

Equilibrate the column with mobile phase with the chromatographic conditions for stable baseline. Inject blank and record the chromatogram. Inject standard preparation in five replicates and record the chromatograms. It should comply with the system suitability criteria as mentioned.





#### FINISHED PRODUCT SPECIFICATION AND TEST METHOD

NAME OF PRODUCT: TENELIGLIPTIN TABLETS 20 MG

PAGE NO .:- Page 5 of 5

- Theoretical plate for Teneligliptin peak should not be less than 2000
- The relative standard deviation for area of Teneligliptin peak should not be more than 2.0 %
- The tailing factor for Teneligliptin should not be more than 2.0.

#### 13.5 Procedure:

Inject sample preparation in duplicate and record the chromatogram. Inhibit the integration due to blank peak in the chromatogram of sample preparation.

#### Calculations:

mg/tab of Teneligliptin = 
$$\frac{At}{As} \times \frac{Ws}{100} \times \frac{100}{Wt} \times \frac{P}{100} \times Aw \times \frac{426.57}{628.86} \times \frac{100}{LC}$$

Where,

A<sub>T</sub> = Area of the peak due to Teneligliptin obtained in the chromatogram of sample preparation

As = Mean area of the peak due to Teneligliptin obtained in the chromatogram of standard preparation.

Ws = Weight of Teneligliptin hydrobromide hydatre working standard taken for standard preparation, in mg.

P = Purity of Teneligliptin hydrobromide hydatre working standard, on as is basis.

LC = Label claim in mg.

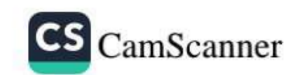
Aw = Average weight in mg.

W<sub>T</sub> = Weight of sample taken for sample preparation, in mg.

426.57 = Molecular weight of Teneligliptin.

628.86 = Molecular weight of Teneligliptin Hydrobromide Hydrate

Limit: Not less than 90.00 % and not more than 110.00 % of the label claim







## FINISHED PRODUCT SPECIFICATION AND TEST METHOD

NAME OF PRODUCT: TENELIGLIPTIN TABLETS 20 MG

PAGE NO .:- Page 2 of 4

### TEST METHOD

1) Description: White coloured, round shaped, film coated tablets, plain on both sides.

#### 2) Identification:

The retention time of the principal peak in the chromatogram of sample preparation should correspond to that of the standard preparation as obtained in the "Assay".

3) Average weight:

TABLETS-1	1:	285 mg.	TABLETS-6	:	287. mg.
TABLETS-2	:	280 mg.	TABLETS-7	:	296. mg.
TABLETS-3	1:	285 mg.	TABLETS-8	1	287 mg.
TABLETS-4	:	292 mg	TABLETS-9	:	286 mg.
TABLETS-5	1:	284 mg.	TABLETS-10	:	282 mg.

AVERAGE WEIGHT:- 286.4 mg .

LIMIT: 283 MG ± 7.5%

#### 4) Uniformity of Weight:

Select randomly 10 tablets and weigh individual tablet. Calculate average, the minimum and maximum value.

TABLETS-1	:	2 86 mg	TABLETS-6	:	290 mg.
TABLETS-2	:	285mg	TABLETS-7	:	285 mg
TABLETS-3	1:	280 mg	TABLETS-8	:	287 mg
TABLETS-4		284 mg	TABLETS-9	:	296 mg
TABLETS-5	:	292 mg	TABLETS-10	:	287 mg

MINIMUM WEIGHT:- 280mg .

LIMIT: 283 MG ±7.5% (BETWEEN 261.8 MG AND 304.2 MG)





## FINISHED PRODUCT SPECIFICATION AND TEST METHOD

NAME OF PRODUCT: TENELIGLIPTIN TABLETS 20 MG

PAGE NO:- Page 4 of 4

## 6) Assay (By HPLC):

## **Chromatographic Conditions:**

Column	1:1	C18 (150 mmx 4.6mm)		
Pump mode	:	Isocoatic		
Mobile Phase	1:	Buffer: Acetonitrik.		
Flow rate	:	1.00 mp min .		
Injection volume	:	20-42.		
Column Temperature	:	300C.		
Wavelength	:	1.5 times of retention time peux.		

### Preparation of solutions:

- Standard preparation:
- Sample preparation:

Standard Weight :- 20 mg.

Sample Weight :- 286-8 mg

Average Weight :- 286.8 mg

Potency :- 99.85 7.

Calculations:

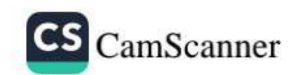
% of Teneligliptin = 
$$\frac{At}{As} \times \frac{Ws}{100} \times \frac{100}{Wt} \times \frac{P}{100} \times Aw \times \frac{426.57}{628.86} \times \frac{100}{LC}$$

1) 
$$\frac{11960800}{11135986} \times \frac{29.43}{100} \times \frac{100}{2868} \times \frac{P}{100} \times 2868 \times \frac{426.57}{628.86} \times \frac{100}{29}$$

2) 
$$\frac{11947512}{11136986} \times \frac{29.43}{100} \times \frac{100}{286.8} \times \frac{P}{100} \times 286.8 \times \frac{426.57}{628.86} \times \frac{100}{20}$$

Average :-

Limit: Not less than 90.00 % and not more than 110.00 % of the label claim







## FINISHED PRODUCT SPECIFICATION AND TEST METHOD

NAME OF PRODUCT: TENELIGLIPTIN TABLETS 20 MG

PAGE NO .:- Page 3 of 4

### 5) Dissolution (By HPLC):

Dissolution Pa	aram	eters :		-	
Medium	1:	Water.	Rotatory Speed	:	75 RPM.
Volume	:	900ml	Temperature	:	37°c.
Apparatus		peddle.(US12-2)	Time	:	45 min .

### Standard Weight :-

Potency:-

Calculations: Teneligliptin (% Drug Release) = 
$$\frac{At}{As} \times \frac{Ws}{100} \times \frac{900}{LC} \times \frac{P}{100} \times \frac{426.57}{628.86} \times 100$$

Tablet 1= 
$$\frac{1.9290}{1.4473} \times \frac{32.43\times5}{100\ 50} \times \frac{900}{20} \times \frac{94.85}{100} \times \frac{426.57}{628.86} \times 100 = 131.73$$

Tablet 2= 
$$\frac{1.8674}{1.4473} \times \frac{32.43 \times 5}{100 50} \times \frac{900}{20} \times \frac{99.85}{100} \times \frac{426.57}{628.86} \times 100 = 127.53$$

Tablet 3= 
$$\frac{1.8199}{1.4473} \times \frac{32.43 \times 5}{10050} \times \frac{900}{20} \times \frac{99.85}{100} \times \frac{426.57}{628.86} \times 100 = 124.32$$

Tablet 4= 
$$\frac{1.7653}{1.4473} \times \frac{3243 \times 5}{10050} \times \frac{900}{20} \times \frac{99.85}{100} \times \frac{426.57}{628.86} \times 100 = 118.48$$

Tablet 5= 
$$\frac{1.7349}{1.4473} \times \frac{3243 \times 5}{10050} \times \frac{900}{20} \times \frac{99.85}{100} \times \frac{426.57}{628.86} \times 100 = 118.48$$

Tablet 6= 
$$\frac{1.8132}{1.4473} \times \frac{32.4385}{100.50} \times \frac{900}{20} \times \frac{99.85}{100} \times \frac{426.57}{628.86} \times 100 = 123.83$$

#### Average:-

Limits: Not less than 80.00 % of labeled amount is dissolved in 45 minutes



	ARNI ANALYTICAI	LS
TITLE	HPLC DATA SHEET	100
Instrument Name :-	High performance liquid charmatography.	Page No
Instrument Make :-	SHTMANZIE	Water Town
Instrument Model No. :-	LC 240 CHT.	
Instrument ID : -	ARNI / INS - OOT	1 of 3
Name Of Student :-	Sanais Baurath Laman.	

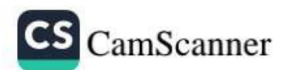
## HPLC DATA SHEET

- HPLC Parameter settings:
  - Make a purging of the mobile phase of all ports & injection port to remove the air bubble from the line.
  - Create a new method by using below parameter.
  - · Save the Method Parameters with a file name.
  - Download the method to the instruments.

#### CHROMATOGRAPHIC PARAMETERS-1

INSTRUMENT PARAMETER	Set Parameters	
Data Aquisitation Time	:	13min.
Pump	:	0.80 m/min.
Port	:	c
Detector (Wavelength)	:	222 nm.
Column Oven Temperature		4000
Degasser		OFF
Autosampler Temperature	:	₹°C.

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	ARNI ANALYTICA	LS	
TITLE	HPLC DATA SHEET		
Instrument Name :-	High Performance Liquid Chromatography		
Instrument Make :-	SHTMODZU	Page No	
Instrument Model No. :-	LC 2do CHI.	1 of 1	
Instrument ID : -	ARNT/IN 5-001		

#### **DATA SHEET**

## NAME OF TEST: - SYSTEM SUITABILITY

## SYSTEM SUITIBILITY CHECK BY INJECTING 3 REPLICATE INJECTIONS OF CAFFEINCE

## • Chromatographic Conditions:

	A stainless steel column
	Dimentions :-
Column	Length :- 15 cm × Diameter:- 4.6 mm; Particle size:-5μm
	Length :- 150 mm × Diameter:- 4.6 mm; Particle size :-5μm
	Stationary Phase :- Packed with octadecylsilyl (C18) silica gel
Data Aquisitation Time	Minutes
Pump (Flow Rate)	1.00 ml/min
Port	A
Detector (Wavelength)	273nm
Column Oven Temperature	30°C
Degasser	Off
Autosampler Temperature	Off

### MOBILE PHASE PREPARATION:-

Prepare a Mixture of 80 volumes of Water and 20 volumes of Methanol. Mix well.

#### STANDARD PREPARATION :-

Weigh accurately 20mg of Caffeine standard to a 100ml volumetric flask. Add 60ml of HPLC grade water and shake to dissolve completely. Slowly makeup the volume upto the mark. Mix well. Further dilute 5ml of the above solution to 50ml volumetric flask, dilute with water to makeup volume.



	ARNI	ANALYTICA	LS
TITLE	HPLC DATA SHEET	A LINE STATE OF THE STATE OF TH	
Instrument Name :-		10 1 30 EXPT - 7311	Page No
Instrument Make :-			ragerio
Instrument Model No. :-			
Instrument ID : -			1 of 1

NAME OF STUDENT :-

### DATA SHEET

## NAME OF TEST :- SYSTEM SUITABILITY

## SYSTEM SUITIBILITY CHECK BY INJECTING 3 REPLICATE INJECTIONS OF CAFFEINCE

· Chromatographic Conditions:

Column	A Stainless steel column.  Diameter: Length = 15cmx 46mm.
Data Aquisitation Time	7 min.
Pump (Flow Rate)	1.00m1/min -
Port	A
Detector (Wavelength)	273 nm
Column Oven Temperature	30°C.
Degasser	off /
Autosampler Temperature	off.

MOBILE PHASE PREPARATION:

prepared a mise of 70 volumes of waver and 20 volumes of Aletonitrile and miseed well.

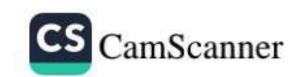
STANDARD PREPARATION :-

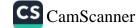
WI. arrandely 20 mg coffine standard to loom! volumenic flask. Add so mi inple grade water and shake it.

SEQUENCE OF INJECTION:-

Name of Solution	No. Of Injection	
Blank		
Standard	2	

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AN	ARNI ANALYTICALS	
TITLE	HPLC DATA SHEET	
Instrument Name :-		
Instrument Make :-		Page No
Instrument Model No. :-		
Instrument ID : -		2 of 3
Name Of Student :-		

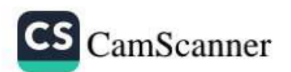
## CHROMATOGRAPHIC PARAMETERS-2

INSTRUMENT PARAMETERS  Data Aquisitation Time :		Set Parameters	
		22min	
Pump		1.20 mitmin	
Port	:	Þ.	
Detector (Wavelength)	1 :	2.60 nm	
Column Oven Temperature	:	30%	
Degasser	:	off.	
Autosampler Temperature	:	1500.	

## CHROMATOGRAPHIC PARAMETERS-3

INSTRUMENT PARAMETERS		Set Parameters
Data Aquisitation Time	:	30 min
Pump	:	1.60 mt min.
Port	:	B-
Detector (Wavelength)	:	260 nm
Column Oven Temperature	:	30 °C
Degasser	:	970
Autosampler Temperature	- :	15%

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	ARNI ANALYTICALS	
TITLE	HPLC DATA SHEET	
Instrument Name :-		Page No
Instrument Make :-		
Instrument Model No. :-		Sully and the Var
Instrument ID:-		3 of 3
Name Of Student :-		

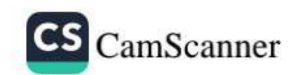
## • CHROMATOGRAPHIC PARAMETERS-4

INSTRUMENT PARAMETERS	198	Set Parameters
Data Aquisitation Time	:	I o min .
Pump	:	1.00ml/min.
Port	:	A.
Detector (Wavelength)	:	216 RM
Column Oven Temperature		3000
Degasser		on.
Autosampler Temperature	:	1000.

## CHROMATOGRAPHIC PARAMETERS-5

INSTRUMENT PARAMETERS		Set Parameters	
Data Aquisitation Time	:	20min	
Pump	:	0 -00 m1/min	
Port	:	A.	
Detector (Wavelength)	:	OFF	
Column Oven Temperature	:	017	
Degasser	:	off	
Autosampler Temperature	:	off.	

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ARNI ANALYTICALS		S
TITLE	TITLE MONTHLY CALIBRATION RECORD OF ANALYTICAL BALANCE	
Instrument Name :-		Page No
Instrument Make :-		
Instrument ID: -		1 of 3

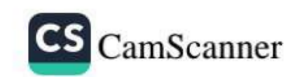
## MONTHLY CALIBRATION RECORD

## Calibration by using Weights:

Observation Table:

Sr. No.	Reference Weight in g	Observed Weight in g	Weight in g (Limit: ± 0.1%)
1	200.0000	199.980	199.8000 to 200.2000
<b>)</b> <sup>2</sup>	100.0000	98.750	99.9000 to 100.1000
3	50.0000	50. 714	49.9500 to 50.0500
4	20.0000	19.055	19.9800 to 20.0200
5	10.0000	10.002	9.9900 to 10.0100
6	5.0000	5.083	4.9950 to 5.0050
7	2.0000	1.329	1.9980 to 2.0020
8	1.0000	0.925	0.9990 to 1.0010
9	0.5000	0.152	0.4995 to 0.5005
10	0.2000	0.105	0.1998 to 0.2002
. 11	0.1000	0.058	0.0999 to 0.1001
12	0.0500	0.038	0.0499 to 0.0501
13	0.0200	0.014	0.0199 to 0.0200
14	0.0100	0.013	0.0099 to 0.0100
15	0.0050	0.008 .	0.0049 to 0.0051

Conclusion: The observed weights are within limit/ out-of-limit.





	ARNI ANALYTICALS		
TITLE	MONTHLY CALIBRATION RECORD OF ANALYTICAL BA	LANCE	
Instrument Name :-		Page No	
Instrument Make :-			
Instrument ID:-		2 of 3	

## 2. Test for Linearity:

Sr. No.	Selected Weights in g	Observed Weight in g
1	50	50.714
2	100	98.790
3	200	200.067.

Conclusion: The observed weights are Consistent/not Consistent.

ANALYSED BY

CHECKED BY

## 3. Test for Eccentricity:

1 2 A 3 4

Sr. No.	Weight Observed in g		Differen	ce in g	Limit
1.	At Centre- (A)	50 - 714			
2.	At Corner 1 (B)	50.719	B-A =	-0.004	
3.	At Corner 2 (C)	50.710	C-A =	- 0.004	± 0.1 %
4.	At Corner 3 (D)	50.708	D-A =	- 0.006	
5.	At Corner 4 (E)	50.709	E-A =	- 0.005	

Conclusion: The maximal Differential Eccentricity error is within limit/out of limit of Std. deviation.

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AN	ARNI ANALYTICALS		
TITLE	MONTHLY CALIBRATION RECORD OF ANALYTICAL BALANCE		
Instrument Name :-		D M.	
Instrument Make :-		Page No	
Instrument ID:-		3 of 3	

## 4. Test for Repeatability:

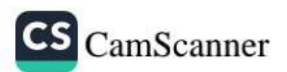
Selected Weight in g: 100 9

Sr. No.	Observed Weight in g	Sr. No.	Observed Weight in g	Limit
1	98.790	6	98.796	
2	98 · 792	7	98.793	
3	98.794	8	98.794	± 0.1 %
4	98 . 795	9	98.794	
5	98 . 793	10	98.794.	

Conclusion: Individual measurement deviation from average value exceeds/ does not exceed standard deviation.

Remark: The instrument is found Satisfactory/ unsatisfactory for its use.

ANALYSED BY





M	ARNI ANALYTICALS		
TITLE MONTHLY CALIBRATION RECORD OF ANALYTICAL BALANCE			
Instrument Name :-	Analytical Balance.	Page No.	
Instrument Make :-	wenser.	Tage No.	
Instrument Model No.:-	DS 8000	1 of 3	
Instrument ID:-			

## NAME OF STUDENT:-

## MONTHLY CALIBRATION RECORD

1. Calibration by using Standard certified weights:

		-		
Observa	tion	Ta	bl	e:

Sr. No.	Reference Weight in g	Observed Weight in g	Weight in g (Limit: ± 0.1%)
1	200.0000	200.022.	199.8000 to 200.2000
2	100.0000	98.770.	99.9000 to 100.1000
3	50.0000	50.699.	49.9500 to 50.0500
4	20.0000	19.051.	19.9800 to 20.0200
5	10.0000	9.997-	9.9900 to 10.0100
6	5.0000	5.080.	4.9950 to 5.0050
7	2.0000	1.230.	1.9980 to 2.0020
8	1.0000	0.923.	0.9990 to 1.0010
9	0.5000	0.191	0.4995 to 0.5005
10	0.2000	0.106	0.1998 to 0.2002
11	0.1000	0.057.	0.0999 to 0.1001
12	0.0500	0.035	0.0499 to 0.0501
13	0.0200	0.014	0.0199 to 0.0200
14	0.0100	0.008.	0.0099 to 0.0100
15	0.0050	0.005.	0.0049 to 0.0051

Conclusion: The observed weights are within limit/ out of limit.

ANALYSED BY





AN	ARNI ANALYTICALS		
TITLE MONTHLY CALIBRATION RECORD OF ANALYTICAL BALANCI		TICAL BALANCE	
Instrument Name :-			
Instrument Make :-		Page No.	
Instrument Model No. :-		2 of 3	
Instrument ID:-		2013	

## 2. Test for Linearity:

Sr. No.	Selected Weights in g	Observed Weight in g
1	5	5.081.
2	10.	10.002.
3	20.	18.375.

Conclusion: The observed weights are Consistent/not Consistent.

## ANALYSED BY

CHECKED BY:-

3. Test for Eccentricity:

1 2 A 3 4

Sr. No.	Weight Observed in g	Difference in g	Limit
1.	At Centre- (A)   0.000.		11
2.	At Corner 1 (B) g.ggg	B-A =	
3.	At Corner 2 (C) q.998.	C-A =	± 0.1 %
4.	At Corner 3 (D) 10.000	D-A =	
5.	At Corner 4 (E) 9.997.	E-A =	

Conclusion: The maximal Differential Eccentricity error is within limit/out of limit of Std. deviation.

ANALYSED BY





AN	ARNI ANALYTICALS		
TITLE	MONTHLY CALIBRATION RECORD OF ANALYTI	ICAL BALANCE	
Instrument Name :-		D M.	
Instrument Make :-		Page No.	
Instrument Model No. :-		3 of 3	
Instrument ID : -			

## 4. Test for Repeatability:

Selected Weight in g: 10 9m.

Sr. No.	Observed Weight in g	Sr. No.	Observed Weight in g	Limit
1	9.997.	6	9.998	
2	9.998-	7	9.996.	
3	10.000 .	8	9.997.	± 0.1 %
4	9.999.	9	9.997.	
5	9.998.	10	9.998.	

Conclusion: Individual measurement deviation from average value exceeds/ does not exceed standard deviation.

Remark: The instrument is found Satisfactory/ unsatisfactory for its use.

ANALYSED BY





17/	ARNI ANALYTICA	LS
TITLE	DAILY CALIBRATION RECORD OF pH-METER	
Instrument Name :-	PH-Meter.	Page No
Instrument Make :-	LABMAN	1 of 1
Instrument Model No. :-		-
Instrument IL: -	ARNI/INS-005	

## DAILY CALIBATION RECORD

- Procedure: Refer SOP No.: SOP/ARN/INS-005
- Preparation Of Solutions:
- pH-4.01:-
  - Transfer the capsule content in a 100ml volumetric flask using a funnel.
  - Dissolve the contents in 10 ml of distilled water and then make it up to 100 ml with distilled water.
  - This solution will have a pH of  $4.0 \pm 0.05$  at  $25^{\circ}$ C.
- pH-7.00:-
  - Transfer the capsule content in a 100ml volumetric flask using a funnel.
  - Dissolve the contents in 10 ml of distilled water and then make it up to 100 ml with distilled water.
  - This solution will have a pH of 7.0 ±0.05 at 25°C.
- pH-9.20:-
- Transfer the capsule content in a 100ml volumetric flask using a funnel.
  - Dissolve the contents in 10 ml of distilled water and then make it up to 100 ml with distilled water.
  - This solution will have a pH of 9.20 ±0.05 at 25°C.
- Observation Table:

Table:		pН	
Sr. No.	Date	4.00 (± 0.05)	7.00 (± 0.05)
1.	16/12/2022.	3.96	6.87.

Slope = 95 %.



	ARNI ANALYTICALS	
TITLE	DAILY CALIBRATION RECORD OF pH-METER	
Instrument Name :-		Page No
Instrument Make :-		A CONTRACTOR OF THE PARTY OF TH
Instrument Model No. :-		1 of 1
Instrument ID:-		

## DAILY CALIBATION RECORD

- Procedure: Refer SOP No.: SOP/ARN/INS-005
- Preparation Of Solutions:
- pH-4.01 :-
  - Transfer the capsule content in a 100ml volumetric flask using a funnel.
  - Dissolve the contents in 10 ml of distilled water and then make it up to 100 ml with distilled water.
  - This solution will have a pH of 4.0 ±0.05 at 25°C.
- pH-7.00 :-
  - Transfer the capsule content in a 100ml volumetric flask using a funnel.
  - Dissolve the contents in 10 ml of distilled water and then make it up to 100 ml with distilled water.
  - This solution will have a pH of 7.0 ±0.05 at 25°C.
- pH-9.20 :-
  - Transfer the capsule content in a 100ml volumetric flask using a funnel.
  - Dissolve the contents in 10 ml of distilled water and then make it up to 100 ml with distilled water.
  - This solution will have a pH of 9.20 ±0.05 at 25°C.

#### 1 • Observation Table:

Sr. No.		pН	the Dan Land
	Date	4.00 (± 0.05)	7.00 (± 0.05)
1.	10/12/2017	3.99	6.86

Slope : 97 %

PERFORMED BY





ARNIANALYTICALS		LS
TITLE	DISSOLUTION TEST APPARATUS WORKSHEET	
Instrument Name :-	Dissolution test apprulus.	Page No.
Instrument ID : -	ARNI / TNS -003	
Instrument Model No. 1- Db - 8000 .		1 of 1
Name Of Students	Sanap Saugabh Lamman.	

## NAME OF TEST :-

TRIAL FOR DISSOLUTION TEST.

### **DISSOLUTION CONDITIONS:-**

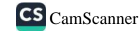
Dissolution Media	WATER	
Media Volume	900 mL	
Apparatus	USP TYPE II PADDLE	
RPM	100	
Temperature	37.0 ± 0.5°C	
Time	45 Minutes	

## PREPARATIONS:-

Pour 900 mL of dissolution medium in each vessel. Allow sufficient time for the dissolution medium to equilibrate at 37°C ± 0.5°C. Adjust stirring element speed to 100 rpm. Place one capsule in each of six paddle and adjust the paddle in the dissolution medium so that there is a distance of 25 mm ± 2 mm between the bottom of the paddle and inside bottom of the vessel. Start the apparatus. At the end of specified time interval, withdraw 10 mL aliquot from a zone midway between the surface of the dissolution medium and at top of the rotating paddle. Further dilute 2ml of the above solution to 25ml with dissolution medium.







	ARNI ANALYTICALS
TITLE DIS	SOLUTION TEST APPARATUS WORKSHEET
Instrument Name :-	
Instrument ID : -	Page I
Instrument Model No. :-	
Name Of Students	1 of

## NAME OF TEST :-

TRIAL FOR DISSOLUTION TEST.

DISSOLUTION CONDITIONS:-

Dissolution Media	Mailor
Media Volume	Water
Apparatus	900m1
57.5	SIGGAS IT STET CISO
RPM	100
Temperature	37.0 t 0.5°C.
Time	45 minitus.

#### PREPARATIONS:-

poured gooms of dissolution medium in each vessle.

17110wed sufficient time for the dissolution medium
to equilibrate at 320c ± 0.5%.

Adjusted Stirring element speed to loss rpm. Placed one capsule in each of six peddle in the dissolution midium so that there is distance of 25mm + 2mm slanded the apparatus. At the end of specified time interval. Withdraweled lamb a liquid. Partner diluted 2ml of the above solution to 25ml with dissolution time.

ANALYSED BY

