Hetero Drugs Limited (Unit-I)

Sy.No.s, 213, 214 & 255, Bonthapally Village, Gummadidala Mandal, Sangareddy District, Telangana state, IN Phone: + 091-8458-275314/275777,Fax:+91-8458-275271

CERTIFICATE OF ANALYSIS

Product

: Nebivolol Hydrochloride

Batch No

: NB17070150 Date of Manufacture : August - 2017

Analytical Report No.: NB0170/17

Reference STP No.

: NB-011

Reference

: In-House

Batch Quantity Date of Analysis : 26.95 Kg : 12/08/2017

Retest Date

: July - 2022

-	1	Status		: Initial certification	
S.No	Test	Specifications	Results	Reference	
1	Description	A white to off white crystalline powder	A white crystalline powder	Visual inspection	
2	Solubility	Sparingly soluble in dimethylformamide, slightly soluble in Methanol, Very slightly soluble in water and practically insoluble in 0.1M HCl.		Visual inspection	
	Identification by a) IR	The infra red absorption spectrum of the finely ground sample in KCl dispersion compressed into a disc should exhibits maxima only at the same wave numbers as that of asimilar preparation of Nebivolol Hydrochloride Working Standard.	Matches with the	Ph.Eur.<2.2.24>	
3	b) HPLC	The retention time of the principal peak obtained in Assay preparation-1 should matches with that of the standard preparation.	Matches with the standard	In-House	
	c) Chlorides test	Should meet the requirement for chloride.	Complies	Ph.Eur.<2.3.1>	
	d)XRD	The diffractogram of the test sample should match with that of Nebivolol HCl working standard.	Matches with the standard diffractogram	Ph.Eur.<2.9.33>	
4	Melting Range	Should be between 225.0°C and 235.0°C	225.8°C to 227.2°C	Ph.Eur.<2.2.14>	
5	Chloride content (on dried basis)	Should be between 7.8 % w/w and 8.5 % w/w	8.0 % w/w	1n-House	
6	Loss on drying	Not more than 1.0 % w/w	0.17 % w/w	Ph.Eur.<2.2.32>	
7	Palladium content	Not be more than 5 ppm	Below QL (QL=0.2 ppm)	In-House	
8	Sulfated ash	Not more than 0.1 % w/w	0.06 % w/w	Ph.Eur.<2.4.14>	
9	Heavy Metals	Not more than 0.002 % w/w	Less than 0.002 % w/w	Ph.Eur.<2.4.8> Method-C	
10	Chromatographic purity byHPLC	[2S*[1R*, 5R* (S*)]]- α , α ' – [iminobis(methylene)bis[6-fluoro-3,4-dihydro-2H-1-benzopyran-2-methanol] Related compound-A : Not more than 0.15 %	0.05% (QL=0.004%)		
		Related compound-A + Related compound -B (from chiral purity by HPLC): Not more than 0.15 %	0.05%	In-House	
		Maximum single unknown impurity: Not more than 0.10 %	0.04%		
		Total impurities : Not more than 0.50 %	0.09%		
	Chiral purity by HPLC	[2S*[1R*, 5S* (S*)]]-α,α' – [iminobis(methylene)bis [6-fluoro-3,4-dihydro-2H-1-benzopyran-2-methanol] Related compound-B:: Not more than 0.15 %	Below QL (QL=0.02%)	In-House	
		Nebivolol D-Isomer: Should be between 48.5 to 51.5 %	49.8%		
		Nebivolol L-Isomer: Should be between 48.5 to 51.5 %	50.2%		

Compiled by:

14/08/1017

Checked by:

Date:

in loglix

Authorised signatory:

Date:

4180141

QC-NB-COA-013-05

Date:

Effective date: 05/06/2017

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CERTIFICATE OF ANALYSIS

Product

: Nebivolol Hydrochloride

Batch No

: NB17070150 Date of Manufacture : August - 2017

Analytical Report No.: NB0170/17

: NB-011-04, NB-005-03

Reference

: In-House

Batch Quantity Date of Analysis

Reference STP No.

: 26.95 Kg : 12/08/2017

Retest Date

Statue

: July - 2022

~ ~ ~		T		Status	: Initial certification
S.No.	Test		Specifications	Results	Reference
12	Residual solvents by GC* (Method-I)	Methanol	: Not more than 1500 ppm	49 ppm (QL=7ppm)	In-House
		Ethanol	: Not more than 1000 ppm	Below QL (QL=13ppm)	
		Di-isopropyl ether	: Not more than 50 ppm	Below QL (QL=8ppm)	
		Ethyl acetate	: Not more than 1000 ppm	Below QL (QL=23ppm)	
		Toluene	: Not more than 400 ppm	Below QL (QL=15ppm)	
	Residual solvents by GC* (Method-II)	Acetic acid	: Not more than 5000 ppm	Below QL (QL=200ppm)	
13	Assay by HPLC (on dried basis)	Should be between 98.5	% w/w and 101.5 % w/w	99.3 % w/w	In-House
14 1	Monomethyl amine content by IC	Should not be more than	100 ppm	31 ppm	In-House
	FOB & B-spot content by HPLC	(±)-[1S*(S*)]-6-Fluoro- Benzopyran (FOB): Shou	3,4dihydro-2-oxiranyl-2H-1- uld be not more than 37 ppm	Below QL (QL=12ppm)	In-House
		(±)-[1S*(R*)]-6-Fluoro- Benzopyran (B-spot): Sh	-3,4dihydro-2-oxiranyl-2H-1- nould be not more than 37 ppm	Below QL (QL=12ppm)	
16	Particle size analysis	d(0.90) should be less that	an 25µm	6.0 μm	In-House

* No potential for the class-1 solvents as specified by ICH to be present in Nebivolol hydrochloride, as they are not used in the manufacturing process. The material if tested for these solvents, will comply with the established standards.

The product conforms to the above Specifications.

Compiled by:

14/08/0017

Checked by:

Date:

14/08/12

Authorised signatory:

Date:

14/08/17

QC-NB-COA-013-05

Date:

Effective date:05/06/2017

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