



Cambrex Profarmaco Milano S.r.L.

Via Curiel 34, 20067 Paullo (MI) - ITALY

Tel.: +39 02 90 62 60.1

Certificate of Analysis

Product	AMIODARONE HYDROCHLORIDE	C.A.S. n.	19774-82-4
Batch	400508	Formula	C25H29I2NO3.HCl
Production date	July 2016	M.W.	681.8
Expiration Date	July 2021	T.S.	016.010
Analysis	July 20 2016		
Coa Number	CA3.009		

DETERMINATION	SPECIFICATION	RESULT
DESCRIPTION	White crystalline powder, odourless	COMPLIES
SOLUBILITY	Methylene chloride	Clear solution COMPLIES
APPEARANCE OF SOLUTION	Methanol	Clear and NMCT GY5 COMPLIES
COLOR	Abs.at 420 nm in Methanol	NMT 0.100 0.050
IDENTIFICATIONS	IR spectrum	Conforms to standard Positive COMPLIES
RESIDUE ON IGNITION		NMT 0.1 % 0.0
HEAVY METALS		NMT 20 ppm COMPLIES
LOSS ON DRYING		NMT 0.5 % 0.0
WATER		NMT 0.5 % 0.1
pH		3.2 / 3.8 3.6
IODIDES		NMT 150 ppm COMPLIES
IMPURITY H BY TLC	Diethylaminoethylchloride HCl (EP/USP-H)	NMT 0.01 % ND
RELATED SUBSTANCES BY HPLC	Diiodo butyl (EP/USP-D)	NMT 0.200 % 0.004
	Mono iodo Amiodarone (EP/USP-C)	NMT 0.100 % ND
	Bis desiodo Amiodarone (EP/USP-A)	NMT 0.100 % ND
	Any unspecified impurity	NMT 0.100 % 0.033
	Total unspecified impurities	NMT 0.300 % 0.063
	Total impurities	NMT 0.500 % 0.067
ASSAY	By titration (on dried basis)	98.5 / 101.0 % 100.7
	Assay by HPLC (on dried basis)	98.5 / 101.0 % 99.4
RESIDUAL SOLVENTS BY GLC	Ethanol	NMT 2000 ppm 303
	Acetone	NMT 50 ppm 2
	Chloroform	NMT 50 ppm ND
	Methylene Chloride	NMT 100 ppm ND

FAGRON IBÉRICA, S.A.U
Corresponde al lote de Fagron:

16K 07/2016

Director Técnico

This batch has been manufactured, packaged and tested in accordance with EU GMP Guideline Volume 4 Part II (ICHQ7)

The product conforms to requirements of:
EuPh8.0 - USP39

Approved by Qualified Person / Quality Director

Roberto Baima

07-21-2016

This Certificate of Analysis has been approved by the Qualified Person / Quality Director and produced automatically with validated electronic signature