



CERTIFICATE OF ANALYSIS

Product **MINOXIDIL-USP35/EP 7.4** Code N° **2610290** Batch N° **1241076**

Manufacturing date **Jan 2013** Retesting date (where applicable) **Dec 2017** Expiration date (where applicable)

Analysis Number		N4PF011203		Limits	
Test	Results	M.U.	Min	Max	
Description	White, crystalline powder		White, crystalline powder	Off-white	
Identification					
- IR	Conforms to Standard		Conforms to Standard		
- UV	Conforms		Conforms		
Loss on drying	0,1	%		0,5	
Residue on ignition	0,05	%		0,1	
Heavy metals	< 20	ppm		20	
Appearance of Solution	Not more coloured than Y-6		Not more coloured than Y-6		
Assay (perchloric acid) :	100,1	%dried	98,5	101,0	
Assay HPLC :	97,3	%dried	97,0	103,0	
Related Substances HPLC :					
- 2,4-Diamino-6-Chloro-Pyrimidine	< 0,05	%		0,2	
- 2,4-Diamino-6-Chloro-Pyrimidine-3-Oxide	< 0,05	%		0,2	
- 2,4-Diamino-6-Piperidinpyrimidine	< 0,05	%		0,2	
- Largest unknown Impurity	< 0,05	%		0,10	
- Total Impurities	< 0,05	%		1,5	
OVI (USP) : Not used			Not tested		
Residual Solvents (GC) :					
- ICH Class 1 Solvents: Not used			Not tested		
- ICH Class1 Solvents : Not used			Not tested		
- ICH Class 3 Solvents:					
-2-propanol	487	ppm	< 24 (LOQ)	2000	
- Acetone	21	ppm	< 12 (LOQ)	1000	
Packaging and Storage		Preserve well-closed container			

This batch has been manufactured, packaged and tested in accordance with EU GMP Guideline Volume 4 Part II (ICH Q7A).
Manufacturing site of the Active Pharmaceutical Ingredient : OLON S.P.A. - GARBAGNATE MIL.se

Date

February 11, 2013

D.C. Manager (Roberto Quaglia)