



Certificate No: IT-API/39/H/2019

CERTIFICATE OF GMP COMPLIANCE OF A MANUFACTURER

Part 1

Issued following an inspection in accordance with Art. 111(5) of Directive 2001/83/EC

The competent authority of Italy confirms the following: The manufacturer INDUSTRIALE CHIMICA S.R.L. Site address Via E. H. Grieg, 13 - 21047 SARONNO (VA)

Is an active substance manufacturer that has been inspected in accordance with Art. 111(1) of Directive 2001/83/EC transposed in the following national legislation: D.L. n. 219 of 24th April 2006 art. 53

From the knowledge gained during inspection of this manufacturer, the latest of which was conducted on 2018/09/21, it is considered that it complies with the Good Manufacturing Practice requirements referred to in the principles of GMP for active substances referred to in Article 47 of Directive 2001/83/EC.

This certificate reflects the status of the manufacturing site at the time of the inspection noted above and should not be relied upon to reflect the compliance status if more than three years have elapsed since the date of that inspection. However, this period of validity may be reduced or extended using regulatory risk management principles by an entry in the Restrictions or Clarifying remarks field. The authenticity of this certificate may be verified in EudraGMDP. If it does not appear, please contact the issuing authority.

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Part 2

Name and address of the site: INDUSTRIALE CHIMICA S.R.L. - Via E. H. Grieg, 13, 21047 SARONNO (VA)

Name of the active Substances manufactured or imported:

11-ALPHA-HYDROXY CANRENONE

ABIRATERONE ACETATE

BIMATOPROST

BRINZOLAMIDE

BUDESONIDE

BUMETANIDE

CANRENONE

DALFAMPRIDINE

DESOGESTREL

DIENOGEST

DOFETILIDE

DROSPIRENONE

EPLERENONE

ESTRADIOL HEMIHYDRATE

ESTRONE

ETHINYLESTRADIOL

ETONOGESTREL

FENOTEROL HYDROBROMIDE

PHENTOLAMINE MESYLATE

FLUTICASONE PROPIONATE

FORMOTEROL FUMARATE DIHYDRATE

FULVESTRANT

GESTODENE

GESTONORONE ACETATE

GUANFACINE HYDROCHLORIDE

IDEBENONE

INDACATEROL MALEATE

INDAPAMIDE

LATANOPROST

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LEVONORGESTREL

LEVONORGESTREL ACETATE

LEVONORGESTREL BUTYRATE

LOTEPREDNOL ETABONATE

MEMANTINE HYDROCHLORIDE

MIDODRINE HYDROCHLORIDE

MOMETASONE FUROATE

MOMETASONE FUROATE MONOHYDRATE

NOMEGESTROL ACETATE

NORGESTIMATE

PRASTERONE

PRASTERONE ACETATE

PREDNISOLONE

SALMETEROL XINAFOATE

SULFAMETHYLTHIAZOLE

TAFLUPROST

TIBOLONE

TIOTROPIUM BROMIDE MONOHYDRATE

TIOTROPIUM BROMIDE SOLVATE

TRAVOPROST

TRIMEGESTONE

ULIPRISTAL

ULIPRISTAL ACETATE

ZOPICLONE

3 Manufacturing Operations - Active Substances

ABIRATERONE ACETATE

3.1	Manufacture of Active Substance by Chemical Synthesis	
	3.1.1.	Manufacture of active substance intermediates
		Special Requirements
		Other: Hormones or substances with hormonal activity
	3.1.2.	Manufacture of crude active substance

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	3.1.3. Salt formation / Purification steps: Crystallisation
3.5	General Finishing Steps
	 3.5.1. Physical processing steps drying, milling/micronisation, sieving 3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot
3.6	numbering) of the active substance)
3.0	Quality Control Testing
	3.6.1. Physical / Chemical testing

BIMATOPROST

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	Special Requirements
	Other:
1	Prostagladins
5)	3.1.2. Manufacture of crude active substance
)	3.1.3. Salt formation / Purification steps:
	Crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps
	Drying
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance)3.5.3. Secondary Packaging (placing the sealed primary package within an

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	outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

BRINZOLAMIDE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
	Crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps
	drying,micronisation, sieving
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a
	packaging material which is in direct contact with the substance)
5	3.5.3. Secondary Packaging (placing the sealed primary package within an
2)	outer packaging material or container. This also includes any labelling of the
	material which could be used for identification or traceability (lot
	numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

3 - Manufacturing Operations - Active Substances

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BUDESONIDE		
3.1	Manufacture of Active Substance by Chemical Synthesis	
	3.1.1. Manufacture of active substance intermediates Special Requirements	
	Other: Hormones or substances with hormonal activity 3.1.2. Manufacture of crude active substance	
	3.1.3. Salt formation / Purification steps:	
	Crystallisation	
3.5	General Finishing Steps	
	 3.5.1. Physical processing steps drying, micronisation, sieving 3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot 	
	numbering) of the active substance)	
3.6	Quality Control Testing	
	3.6.1. Physical / Chemical testing	

BUMETANIDE

3.1	Manufacture of Active Substance by Chemical Synthesis
	 3.1.1. Manufacture of active substance intermediates 3.1.2. Manufacture of crude active substance 3.1.3. Salt formation / Purification steps: Crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps

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3.6	Quality Control Testing 3.6.1. Physical / Chemical testing
	packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot numbering) of the active substance)
	drying,micronisation, sieving 3.5.2. Primary Packaging (enclosing / sealing the active substance within a

DALFAMPRIDINE

3.1	Manufacture of Active Substance by Chemical Synthesis	
	3.1.1. Manufacture of active substance intermediates	
	3.1.2. Manufacture of crude active substance	
	3.1.3. Salt formation / Purification steps:	
	crystallisation	
3.5	General Finishing Steps	
TAL	 3.5.1. Physical processing steps drying, micronisation, sieving 3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot numbering) of the active substance) 	
3.6	Quality Control Testing	
	3.6.1. Physical / Chemical testing	

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DESOGESTREL

3.1	Manufacture of Active Substance by Chemical Synthesis	
	3.1.1. Manufacture of active substance intermediates	
	Special Requirements	
	Other: Hormones or substances with hormonal activity	
	3.1.2. Manufacture of crude active substance	
	3.1.3. Salt formation / Purification steps:	
	Crystallizzazione	
3.5	General Finishing Steps	
	3.5.1. Physical processing steps	
	drying, milling/micronisation, sieving	
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a	
	packaging material which is in direct contact with the substance)	
	3.5.3. Secondary Packaging (placing the sealed primary package within an	
	outer packaging material or container. This also includes any labelling of the	
	material which could be used for identification or traceability (lot	
	numbering) of the active substance)	
3.6	Quality Control Testing	
	3.6.1. Physical / Chemical testing	

3 - Manufacturing Operations - Active Substances

DIENOGEST

3.1	Manufacture of Active Substance by Chemical Synthesis		
		Manufacture of active substance intermediates	
		Special Requirements	

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	Other: Hormones or substances with hormonal activity 3.1.2. Manufacture of crude active substance 3.1.3. Salt formation / Purification steps: Crystallisation
3.5	General Finishing Steps
	 3.5.1. Physical processing steps drying, micronisation, sieving 3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an
	outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

DOFETILIDE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
	crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps drying, micronisation
	 3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the

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	material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

DROSPIRENONE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	Special Requirements
	Other: Hormones or substances with hormonal activity
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
	crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps
	drying, micronisation, sieving
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a
	packaging material which is in direct contact with the substance)
	3.5.3. Secondary Packaging (placing the sealed primary package within an
	outer packaging material or container. This also includes any labelling of the
	material which could be used for identification or traceability (lot
12	numbering) of the active substance)
3.6	Quality Control Testing
15-1	3.6.1.Physical / Chemical testing

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3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	Special Requirements
	Other: Hormones or substances with hormonal activity
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
	Crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps
	drying, micronisation
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance)
	3.5.3. Secondary Packaging (placing the sealed primary package within an
	outer packaging material or container. This also includes any labelling of the
	material which could be used for identification or traceability (lot
	numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

ETHINYLESTRADIOL

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	Special Requirements
	Other: Hormones or substances with hormonal activity
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:

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	crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps drying,
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance)
	3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot
	numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

ETONOGESTREL

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	Special Requirements
,	Other: Hormones or substances with hormonal activity
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
	crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps
	drying, micronisation,
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a
	packaging material which is in direct contact with the substance)
	3.5.3. Secondary Packaging (placing the sealed primary package within an
	outer packaging material or container. This also includes any labelling of the
	material which could be used for identification or traceability (lot

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	numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

FENOTEROL HYDROBROMIDE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
	crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps
	drying, sieving
8	3.5.2. Primary Packaging (enclosing / sealing the active substance within a
Z .	packaging material which is in direct contact with the substance)
I'd	3.5.3. Secondary Packaging (placing the sealed primary package within an
	outer packaging material or container. This also includes any labelling of the
	material which could be used for identification or traceability (lot
	numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

3 - Manufacturing Operations - Active Substances

PHENTOLAMINE MESYLATE

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3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
	crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps drying, sieving
	 3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

FLUTICASONE PROPIONATE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates Special Requirements Other: Hormones or substances with hormonal activity
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps: crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps drying, micronisation, sieving3.5.2. Primary Packaging (enclosing / sealing the active substance within a

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	packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

FORMOTEROL FUMARATE DIHYDRATE

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3.1.2. 3.1.3. Gener 3.5.1. 3.5.2. packag 3.5.3. outer materi numbe Qualit 3.6.1.

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FULVEST	JLVESTRANT	
3.1	Manufacture of Active Substance by Chemical Synthesis	
	3.1.1. Manufacture of active substance intermediates Special Requirements	
	Other: Hormones or substances with hormonal activity	
	3.1.2. Manufacture of crude active substance	
	3.1.3. Salt formation / Purification steps:	
3.5	crystallisation	
3.3	General Finishing Steps	
	3.5.1. Physical processing steps	
	drying, sieving	
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a	
	packaging material which is in direct contact with the substance)	
	3.5.3. Secondary Packaging (placing the sealed primary package within an	
	outer packaging material or container. This also includes any labelling of the	
	material which could be used for identification or traceability (lot	
	numbering) of the active substance)	
3.6	Quality Control Testing	
	3.6.1. Physical / Chemical testing	

GESTODENE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates Special Requirements
	Other: Hormones or substances with hormonal activity 3.1.2. Manufacture of crude active substance

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	3.1.3. Salt formation / Purification steps: crystallisation
3.5	General Finishing Steps
	 3.5.1. Physical processing steps drying, micronisation, sieving 3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

GUANFACINE HYDROCHLORIDE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
	crystallisation
3.5	General Finishing Steps
 3.5.1. Physical processing steps drying, micronisation, sieving 3.5.2. Primary Packaging (enclosing / sealing the active sealing material which is in direct contact with the subsection of the sealed primary processing material or container. This also includes a material which could be used for identification or traceable. 	
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot numbering) of the active substance)

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3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

IDEBENONE

3.1	Manufacture of Active Substance by Chemical Synthesis
7.	3.1.1. Manufacture of active substance intermediates
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
	crystallisation
3.5	General Finishing Steps
packaging material which is in direct contact with the sealed print as a secondary Packaging (placing the sealed print packaging).	
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance)
	3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the
	material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

3 - Manufacturing Operations - Active Substances

INDACATEROL MALEATE

3.1 Manufacture of Active Substance by Chemical Synthesis

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	3.1.1. Manufacture of active substance intermediates 3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps: crystallisation
3.5	General Finishing Steps
	 3.5.1. Physical processing steps drying, micronisation 3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

INDAPAMIDE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
	crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps
	drying, micronisation, sieving
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance)
	3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the

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	material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

LATANOPROST

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	Special Requirements
	Other:
	prostaglandines
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
	Purification
3.5	General Finishing Steps
	3.5.1. Physical processing steps
3	filtration
151	3.5.2. Primary Packaging (enclosing / sealing the active substance within a
	packaging material which is in direct contact with the substance)
3.5.3. Secondary Pack outer packaging materi	3.5.3. Secondary Packaging (placing the sealed primary package within an
	outer packaging material or container. This also includes any labelling of the
	material which could be used for identification or traceability (lot
	numbering) of the active substance)
3.6	Quality Control Testing
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LEVONORGESTREL

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates Special Requirements Other: Hormones or substances with hormonal activity
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps: crystallisation
3.5	General Finishing Steps
	 3.5.1. Physical processing steps drying, milling, micronisation, sieving 3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
2	3.6.1. Physical / Chemical testing

3 - Manufacturing Operations - Active Substances

LEVONORGESTREL BUTYRATE

3.1	Manuf	acture of Active Substance by Chemical Synthesis
	3.1.1.	Manufacture of active substance intermediates Special Requirements
	3.1.2.	Other: Hormones or substances with hormonal activity Manufacture of crude active substance

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	3.1.3. Salt formation / Purification steps: crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps drying, sieving
	 3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the
	material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

LOTEPREDNOL ETABONATE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
A)	Special Requirements
=/	Other: Hormones or substances with hormonal activity
/	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
	crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps
	drying, micronisation, sieving
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance)
	3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the

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	materialwhich could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

MEMANTINE HYDROCHLORIDE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
	crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps drying, micronisation, sieving
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance)
i i	3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the
	material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

3 - Manufacturing Operations - Active Substances

MIDODRINE HYDROCHLORIDE

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3.1/	Manufacture of Active Substance by Chemical Synthesis	
	3.1.1. Manufacture of active substance intermediates	
	3.1.2. Manufacture of crude active substance	
80	3.1.3. Salt formation / Purification steps:	
	crystallisation	
3.5	General Finishing Steps	
	3.5.1. Physical processing steps	
	drying, milling, micronisation, sieving	
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a	
	packaging material which is in direct contact with the substance)	
	3.5.3. Secondary Packaging (placing the sealed primary package within an	
F - 7	outer packaging material or container. This also includes any labelling of the	
	material which could be used for identification or traceability (lot	
	numbering) of the active substance)	
3.6	Quality Control Testing	
	3.6.1. Physical / Chemical testing	

MOMETASONE FUROATE

3.1	Manufacture of Active Substance by Chemical Synthesis
JE,	3.1.1. Manufacture of active substance intermediates Special Requirements Other: Hormones or substances with hormonal activity
	3.1.2. Manufacture of crude active substance3.1.3. Salt formation / Purification steps: crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps drying, micronisation, sieving

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	3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

MOMETASONE FUROATE MONOHYDRATE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	Special Requirements
	Other: Hormones or substances with hormonal activity
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
2	crystallisation
3.5	General Finishing Steps
1/2	3.5.1. Physical processing steps
1/2	drying, micronisation, sieving
0	3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance)
	3.5.3. Secondary Packaging (placing the sealed primary package within an
	outer packaging material or container. This also includes any labelling of the
	material which could be used for identification or traceability (lot
	numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

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NOMEGESTROL ACETATE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	Special Requirements
	Other: Hormones or substances with hormonal activity
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
	cristallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps
	drying, micronisation, sieving
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a
1 2 7 1 9	packaging material which is in direct contact with the substance)
	3.5.3. Secondary Packaging (placing the sealed primary package within an
	outer packaging material or container. This also includes any labelling of the
	material which could be used for identification or traceability (lot
	numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

3 - Manufacturing Operations - Active Substances

NORGESTIMATE

3.1	Manufacture of Active Substance by Chemical Synthesis	
	3.1.1. Manufacture of active substance intermediates	

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	Special Requirements Other: Hormones or substances with hormonal activity 3.1.2. Manufacture of crude active substance 3.1.3. Salt formation / Purification steps:
3.5	crystallisation General Finishing Steps
	 3.5.1. Physical processing steps drying, micronisation, sieving 3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

SALMETEROL XINAFOATE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
	crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps
	drying,micronisation, sieving
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance)
	3.5.3. Secondary Packaging (placing the sealed primary package within an

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	outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

SULFAMETHYLTHIAZOLE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
	crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps
	drying, micronisation, sieving
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a
	packaging material which is in direct contact with the substance)
A	3.5.3. Secondary Packaging (placing the sealed primary package within an
	outer packaging material or container. This also includes any labelling of the
2	material which could be used for identification or traceability (lot
	numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

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TAFLUPF	TAFLUPROST	
3.1	Manufacture ofActive Substance by Chemical Synthesis	
	3.1.1. Manufacture of active substance intermediates Special Requirements Other: prostaglandins 3.1.2. Manufacture of crude active substance 3.1.3. Salt formation / Purification steps:	
3.5	purification General Finishing Steps	
	 3.5.1. Physical processing steps drying 3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot numbering) of the active substance) 	
3.6	Quality Control Testing	
	3.6.1. Physical / Chemical testing	

TIBOLONE

3.1	Manuf	acture of Active Substance by Chemical Synthesis
		Manufacture of active substance intermediates
		Special Requirements
		Other: Hormones or substances with hormonal activity
	3.1.2.	Manufacture of crude active substance
		Salt formation / Purification steps:

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	crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps
	drying, micronisation, sieving
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance)
	3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

TIOTROPIUM BROMIDE MONOHYDRATE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	3.1.2. Manufacture of crude active substance
121	3.1.3. Salt formation / Purification steps:
Jan .	crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps drying
	 3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the
	material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing

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3.6.1. Physical / Chemical testing

3 - Manufacturing Operations - Active Substances

TIOTROPIUM BROMIDE SOLVATE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
	crystallisation
3.5	General Finishing Steps
	 3.5.1. Physical processing steps drying 3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

3 - Manufacturing Operations - Active Substances

TRAVOPROST

3.1	Manufacture of Active Substance by Chemical Synthesis	
	3.1.1. Manufacture of active substance intermediates	

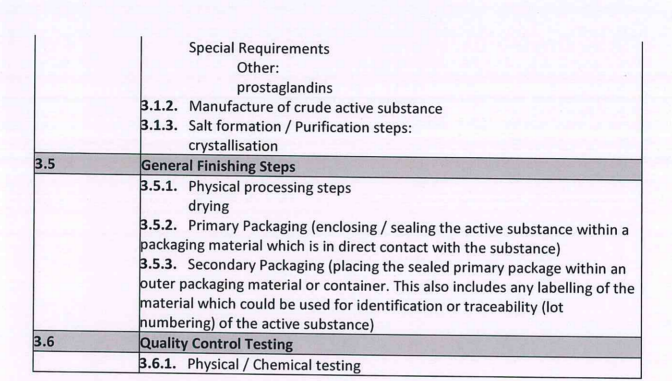
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TRIMEGESTONE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates Special Requirements
	Other: Hormones or substances with hormonal activity 3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps: crystallisation
3.5	General Finishing Steps
	3.5.1. Physical processing steps drying

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	3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance)
	3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

ULIPRISTAL ACETATE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates Special Requirements Other: Hormones or substances with hormonal activity
	3.1.2. Manufacture of crude active substance3.1.3. Salt formation / Purification steps:crystallisation
3.5	General Finishing Steps
	 3.5.1. Physical processing steps drying, micronisation 3.5.2. Primary Packaging (enclosing / sealing the active substance within a packaging material which is in direct contact with the substance) 3.5.3. Secondary Packaging (placing the sealed primary package within an outer packaging material or container. This also includes any labelling of the material which could be used for identification or traceability (lot numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

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ZOPICLONE

3.1	Manufacture of Active Substance by Chemical Synthesis
	3.1.1. Manufacture of active substance intermediates
	3.1.2. Manufacture of crude active substance
	3.1.3. Salt formation / Purification steps:
	crystallisatiion
3.5	General Finishing Steps
	3.5.1. Physical processing steps
	drying, micronisation, sieving
	3.5.2. Primary Packaging (enclosing / sealing the active substance within a
	packaging material which is in direct contact with the substance)
	3.5.3. Secondary Packaging (placing the sealed primary package within an
	outer packaging material or container. This also includes any labelling of the
	material which could be used for identification or traceability (lot
	numbering) of the active substance)
3.6	Quality Control Testing
	3.6.1. Physical / Chemical testing

4. Other Activities - Active Substance:

Importation of:

11-ALPHA-HYDROXY CANRENONE (Confidential); CANRENONE (Confidential); ESTRADIOL HEMIHYDRATE (Confidential); ESTRONE (Confidential); GESTONORONE ACETATE (Confidential); LEVONORGESTREL ACETATE (Confidential); MEMANTINE HYDROCHLORIDE; PRASTERONE (Confidential); PRASTERONE ACETATE (Confidential); PREDNISOLONE (Confidential); ULIPRISTAL (Confidential)

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Restrictions or clarifying remarks:

The Inspectorate adopted a risk-based approach for planning of inspections, therefore the validity of the GMP certificate for this manufacturing site is not more than 42 months from the last general GMP inspection, which was conducted on 2018/09/21. It will still be AIFA's right to re-evaluate the validity of the GMP certificate based on risk profile changes. Imported active substances marked as confidential undergo further processing within the importing site.

Rome, 2019/02/26

Name and signature of the authorised person of the Competent Authority of Republic of Italy

Dott.ssa Marisa Delbò

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