

CERTIFICATE OF ANALYSIS

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|-------------------|---|---------------------|--|
| Report issued to | NEOLPHARMA Grupo Farmaceutico | Report No. | CMC/LAB/NPMP26/2025 |
| Sample Name | MASTERON 100 (Drostanolone Prop.) | ULR No. | NA |
| Manufacturer | NEOLPHARMA Grupo Farmaceutico (Mexico) | Reporting Date | 17-09-2025 |
| Active Ingredient | Drostanolone Propionate | Sample Submitted By | Ing. Ricardo Alvarez Mendez, QA Manager (NEOLPHARMA) |
| Package | 10 ml vial | Exp. Date | 01-2029 |
| Batch / Lot No. | NPMP26 | Sample Qty. | 1 vial |
| Received Date | 15-09-2025 | Mfg. Lic. No. | COFEPRIS 153300201X0092 |
| Sample Condition | Received in sealed intact condition | Ref. No. | NIL |
| Storage Condition | Store below 30C away from direct sunlight | Country of Origin | Mexico |

| | | | |
|--------------------------------|-------------------------------|------------------------|------------|
| Reference to Protocol / Method | USP 2024 . CMC-LAB/SOP/NPTE23 | Analysis Started on: | 15-09-2025 |
| Analysis Completed on: | 17-09-2025 | Review / Release Date: | 18-09-2025 |

RESULT OF ANALYSIS

| S.No | Test Parameter | Claim | Units | Results | Specifications | Test Method |
|------|----------------------------------|-------|-------|----------|--|--------------------|
| 1 | Description | - | - | Complies | Clear pale yellow oily solution, practically free from visible particulates. | Visual Evaluation. |
| 2 | Nominal Volume | 10.0 | ml | 10.05 | 9.5 ml to 10.5 ml | USP <1> |
| 3 | Identification | - | - | Complies | Positive for Drostanolone Propionate | HPLC |
| 4 | Assay of Drostanolone Propionate | 100 | mg/ml | 99.8 | 95.00 - 105.00 mg/ml | HPLC |
| 5 | Sterility | - | - | Complies | No evidence of microbial growth | USP <71> |
| 6 | Bacterial Endotoxin | - | EU/ml | < 0.25 | NMT 1.0 EU/ml | USP <85> |


The sample submitted by NEOLPHARMA Grupo Farmaceutico **COMPLIES** with the declared specifications and is of standard pharmaceutical quality with respect to the parameters tested.

USP = United States Pharmacopela . HPLC = High Performance Liquid Chromatography . KF = Karl Fischer . NMT = Not More Than
 EU = Endotoxin Units . AV = Acceptance Value . COFEPRIS = Comision Federal para la Proteccion contra Risagos Sanitarios


QFB. Laura A. Martinez Rios
 Chief Analyst, Costamed Laboratory




Dr. Arturo Gonzalez Pena
 Technical Director, Costamed Laboratory



**** End of Report ****

Terms and Conditions

1. This report is issued by Costamed Grupo Medico and refers only to the sample tested.
2. Sample will be destroyed one year from the date of issue unless otherwise specified.
3. Total liability of the laboratory is limited to the invoiced amount.
4. All disputes subject to Cosamel, Quintana Roo jurisdiction, Mexico.