

# South African Health Products Regulatory Authority



Licence number: 0000001451.-.1

## LICENCE TO MANUFACTURE MEDICINES (Contract Testing Laboratory for Cannabis Only)

In terms of section 22C(1)(b) of the Medicines and Related Substances Act, 1965

This licence is granted to:

Licence Holder
<b>Ecogreen Analytics (Pty) Ltd</b>
13 Kingfisher Park, Ou Paardeveli Road, The interchange, Somerset West, Western Cape, 7130

### On the following terms and conditions:

The licence holder and the persons described and named in Annexure 1 shall at all times ensure that all medicines manufactured in this facility, irrespective of its registration status, comply with all the provisions of the Medicines and Related Substances Act, 1965, as amended and in particular with sections 14, 18, 18A, 18B, 18C, 19, 20, 22A, 22C, 22G, 33, Regulations 7, 10, 11, 12, 13, 14, 15, 40, 42, 53 and all relevant SAHPRA Guidelines.

This facility is authorised to perform the manufacturing activities depicted in Annexure 1 to this licence.

Boitumalo Senete Makakafeta  
  
17/10/2023 09:44:48(UTC+02:00)

17 October 2023

### CHIEF EXECUTIVE OFFICER

**ORIGINAL ISSUE DATE:** 12 October 2023  
**RENEWAL DATE:** N/A  
**AMENDMENT DATE:** N/A  
**EXPIRY DATE:** 12 October 2028



<b>AUTHORISED MANUFACTURING AND MATERIAL HANDLING ACTIVITIES</b>		
<b>1. MANUFACTURING ACTIVITIES</b>	<b>YES</b>	<b>NO</b>
<b>Sterile, Non-biological Manufacture (includes filling, but not cartoning or labelling)</b>		
Large volume parenteral products		NO
Small volume parenteral products		NO
Other sterile dosage forms: N/A		NO
<b>Non-sterile Manufacture</b>		
Tablets		NO
Capsules		NO
Liquids		NO
Semi-solids		NO
Suppositories		NO
Other non-sterile dosage forms: N/A		NO
<b>Biological Manufacture</b>		
Vaccines		NO
Sera and other immunologicals		NO
Blood and other blood products		NO
Other biological products:		NO
<b>Medical Gas Manufacture</b>		NO
<b>Radioactive Medicines Manufacture</b>		NO
<b>Complementary Medicines Manufacture</b>		NO
<b>2. PACKAGING ACTIVITIES</b>		
Packaging of bulk products and labelling		NO
Re-labelling or redressing		NO
Cartoning or secondary packaging		NO
<b>3. TESTING ACTIVITIES</b>		
Analytical: chemical testing of cannabis samples including hemp, extracts and derivatives of cannabis (CBD, THC, CBN, THCA, Delta 9 THC, Delta 8 THC)	YES	
Microbiological		NO
Sterility		NO
Stability		NO
Animal		NO
Other Testing Activities:		NO
<b>4. DISTRIBUTION ACTIVITIES</b>		
Bulk distribution to wholesale pharmacies		NO
Fine distribution to retail pharmacies and others		NO
<b>5. MATERIALS HANDLED OR STORED AT THIS SITE</b>		
Penicillins (Finished Packed Products Only)		NO
Cephalosporins (Finished Packed Products Only)		NO
Hormones (Finished Packed Products Only)		NO
Cytostatics/Cytotoxics (Finished Packed Products Only)		NO
Bulk Pesticides, Herbicides or Rodenticides (Finished Packed Products Only)		NO
Potent Steroids (Finished Packed Products Only)		NO
Other potent, toxic, sensitising or hazardous materials (Finished Packed Products Only)		NO
<b>6. IMPORT</b>		NO
<b>7. EXPORT</b>		NO
Specific Products Exported:		



**8. PARTICULARS OF THE PERSONNEL RESPONSIBLE FOR OPERATION ON THE PREMISES ON BEHALF OF THE LICENCE HOLDER**

Responsible Pharmacist	Head of Production	Quality Control Person
Not mandatory for a testing Laboratory	Not mandatory for a testing Laboratory	Gregory Andrew Ondrejko
-	-	BSC Chemistry

**9. PARTICULARS OF THE NATURAL PERSON RESPONSIBLE TO THE SOUTH AFRICAN HEALTH PRODUCTS REGULATORY AUTHORITY TO ENSURE COMPLIANCE WITH THE MEDICINES AND RELATED SUBSTANCES ACT, 1965**

Responsible Person	Designation	Residential Address
Gregory Andrew Ondrejko	Technical Director	13 Kingfisher Park, Ou Paardeveli Road, The interchange, Somerset West, Western Cape, 7130
BSC Chemistry		

**10. LICENCE SPECIFIC CONDITIONS**

1. The holder of the licence shall conduct all manufacturing, wholesaling or distribution operations in respect of those medicines for which a registration certificate has been obtained, so as to ensure that the medicines shall conform to the standards of quality, strength and purity applicable to them in accordance with the specifications made by the person to whose order they are manufactured, wholesaled or distributed or the specifications under which the medicines are sold or supplied.
2. Medicine for export for which a registration certificate has not been obtained from the SAHPRA may not be exported without the relevant "Certificate of a Pharmaceutical Product" or alternatively a "Licensing Status of a Pharmaceutical Product" issued by the SAHPRA in terms of the WHO Certification Scheme on the Quality of Pharmaceutical Products moving in International Commerce.

**11. GENERAL CONDITIONS**

- The site to comply with cGMP principles.
- The licence is restricted to activities listed only; any additional activities must be approved by SAHPRA prior to implementation, and
- That any critical changes (Refer to S.A. Guideline: Amendments) to the facility be approved by SAHPRA prior to implementation:

[https://www.sahpra.org.za/wp-content/uploads/2022/06/SAHPGL-LIC-03\\_Guideline-on-how-to-amend-a-current-license\\_June-2022\\_v1\\_DN.pdf](https://www.sahpra.org.za/wp-content/uploads/2022/06/SAHPGL-LIC-03_Guideline-on-how-to-amend-a-current-license_June-2022_v1_DN.pdf)

- Effective implementation of corrective actions to be verified during the next inspection.

**12. SITE SPECIFIC CONDITIONS**

- NONE



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**DOCUMENT HISTORY:**

REVISION	REASON FOR AMENDMENT	DATE
1	First issue of Licence	12 October 2023

