# SOUTH AFRICAN HEALTH PRODUCTS REGULATORY AUTHORITY



Licence number: 00001914MD

# LICENCE TO MANUFACTURE MEDICAL DEVICES

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

To act as a Manufacturer, Distributor and Importer

This licence is granted to:

Licence Holder

Pro-Active Medical Supplies CC

Dowerglen

49 Fairway Boulevard

Edenvale 1609

## 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 medical devices distributed, 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, 22H, 23, 26, 28, 33 and the Regulations relating to Medical Devices 2, 3, 4, 5, 6, 13, 14, 17, 18, 19, 20, 21, 22, 23, 24, 25, 27, 28 and all relevant South African Health Products Regulatory Authority Guidelines

#### This licence consists of 4 pages.

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

Boitumelo Semete-Makokotlela

DocuSigned by

CHIEF EXECUTIVE OFFICER

**ORIGINAL DATE OF ISSUE: 16 April 2021** 

**EXPIRY DATE: 16 April 2026** 

AMENDMENT DATE: N/A

# ANNEXURE 1 00001914MD

# **AUTHORISED MANUFACTURING AND MATERIAL HANDLING ACTIVITIES**

1. MANUFACTURING ACTIVITIES	YES	NO
Sterile Medical Device Manufacture (includes primary packing, but not secondary		
packing such as cartoning or labelling)		Nia
Single use		No
Measuring medical devices		No
Non-invasive medical device		No
Invasive medical devices		No
Active medical devices		No
Inactive medical devices		No
Contraceptive medical devices		No
Combination medical devices		No
Other sterile medical devices (as specified):		No
Non-sterile Manufacture		
Measuring medical devices		No
Non-invasive medical devices		No
Invasive medical devices		No
Active medical devices		No
Inactive medical devices		No
Contraceptive medical devices		No
Combination medical devices		No
Other non-sterile medical devices (as specified):	1	No
Manufactu <mark>re o</mark> f <i>In V<mark>itro Devices (IVDs)</mark></i>	1	
Class A IVD		No
Class B IVD		No
Class C IVD		No
Class D IVD		No
End po <mark>int Ste<mark>rilis</mark>ation of Medical Devices</mark>		No
Manuf <mark>acture of R</mark> adioactive Medical Devices		
Servic <mark>ing and Ref</mark> urbishment of Medical Devices	M	No
	V	
2. PACKAGING ACTIVITIES	/ /	
Packaging of bulk product and labelling		No
Re-labe <mark>lling or redre</mark> ssing		No
Cartoning or secondary packaging	Yes	
Assembly or "kits" / procedure packs		No
	1	
3. TESTING ACTIVITIES	1	
Analytical		No
Microbiological		No
Sterility		No
Stability		No
Animal		No
Other Testing Activities (as specified):		No
4. DISTRIBUTION ACTIVITIES		
	Ver	
Distribution to hospitals and retail pharmacies and other clients: Class A	Yes	
Distribution to hospitals and retail pharmacies and other clients: Class B	Yes	
Distribution to hospitals and retail pharmacies and other clients: Class C	Yes	
Distribution to hospitals and retail pharmacies and other clients: Class D	Yes	

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	YES	NO
5. MATERIALS HANDLED OR STORED AT THIS SITE		
Combination medical devices with Penicillins		No
Combination medical devices with Cephalosporins		No
Combination medical devices with (other) Antibiotics (as specified):		No
Combination medical devices with Hormones		
Combination medical devices with Cytostatics/Cytotoxics		
Bulk Pesticides, Herbicides or Rodenticides	H = A	No
Radioactive material or Radioactive medical devices		No
Other potent, toxic, sensitising or hazardous materials (as specified):		No
6. IMPORT		
Import Class A medical device	Yes	
Import Class B medical device	Yes	
Import Class C medical device	Yes	-
Import Class D medical device	Yes	No
Import Class A IVD		No
Import Class B IVD		No
Import Class C IVD		No
Import Class D IVD		No
Import RUO IVDs		No
	1	
7. EXPORT	1	
Export Class A medical device		No
Export Class B medical device		No
Export Class C medical device	1 1	No
Export Class D medical device	1	No
Export Class A IVD	1	No
Export Class B IVD		No
Export Class C IVD	V	No
Export Class D IVD		No
Export RUO IVDs	1	No

#### 00001914MD

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

Au <mark>tho</mark> rised Representative	Manufactu <mark>re</mark> / Imp <mark>ort</mark> / Dist <mark>rib</mark> ution / Expo <mark>rt Control Perso</mark> n	Quality Control Person
George Algernon	George Algernon Ernest Dewart	George Algernon Ernest
Ernest Dewart		Dewart
Matric	Matric	Matric

# 9. PARTICULARS OF THE LICENCE HOLDER CONTACT (Other than the Authorised Representative)

Name	Contact Details	Address
Mr GAE Dewart	Tel: 011 452 8595	Dowerglen
	Cell: 083 378 3727	49 Fairway Boulevard
	Fax: 086 671 7565	Edenvale
11 /1	Email: proactivemed@worldonline.co.za	1609

# 10. LICENCE SPECIFIC CONDITIONS

1. The holder of the licence shall conduct all manufacturing, distribution or wholesaling operations in respect of those medical devices for which a registration certificate has been obtained, so as to ensure that the medical devices shall conform to the standards of quality, safety and performance applicable to them in accordance with the specifications made by the person to whose order they are manufactured, distributed or wholesaled or the specifications under which the medical devices are sold or supplied.

### 11. ADDITIONAL LICENCE SPECIFIC CONDITIONS (IF REQUIRED)

