# Medicines and Healthcare products Regulatory Agency

## MANUFACTURER'S AUTHORISATION

1: Authorisation Number UK MIA(IMP) 53174

2: Name of authorisation holder REACTA BIOTECH LIMITED

REACTA BIOTECH LIMITED T/A REACTA HEALTHCARE, 2

3: Address(es) of manufacturing site(s)

NEWTECH SQUARE, DEESIDE INDUSTRIAL PARK, DEESIDE,

NEW IECH SQUARE, DEESIDE INDUSTRIAL PARK, DEESIDE

CH5 2NT, UNITED KINGDOM

4: Legally registered address of authorisation holder

INDUSTRIAL PARK, DEESIDE, CH5 2NT, UNITED KINGDOM

5: Scope of authorisation and dosage forms ANNEX 1 and/ or ANNEX 2

6: Legal Basis of authorisation

Part 6 of The Medicines for Human Use (Clinical Trials) Regulations

2004 [SI 2004/1031]

Confidential

7: Name of responsible officer of the competent authority of the member state granting the manufacturing authorisation

8: Authorisation Date 10/09/2024

9: Annexes attached Annex 1 and/or Annex 2

### SCOPE OF AUTHORISATION

#### Annex 2

Name and address of the site:

REACTA BIOTECH LIMITED T/A REACTA HEALTHCARE, 2 NEWTECH SQUARE, DEESIDE INDUSTRIAL PARK, DEESIDE, CH5 2NT, UNITED KINGDOM

**Human Investigational Medicinal Products** 

**Authorised Operations** 

MANUFACTURING OPERATIONS (according to part 1)

## Part 1 - MANUFACTURING OPERATIONS

## [ 1.2 ] Non-sterile investigational medicinal products

[ 1.2.1 ] Non-Sterile Products (processing operations for the following dosage forms)

[ 1.2.1.11 ] Semi-solids

## [ 1.5 ] Packaging

[1.5.1] Primary packaging

[ 1.5.1.11 ] Semi-solids

[1.5.2] Secondary packaging

Issue Date: 10 Sep 2024