Medicines and Healthcare products Regulatory Agency

REGISTRATION OF MANUFACTURER, IMPORTER OR DISTRIBUTOR OF ACTIVE SUBSTANCES TO BE USED AS STARTING MATERIALS IN MEDICINAL PRODUCTS FOR HUMAN USE

Registrant Details

UK API 27830

EUROFINS SELCIA LIMITED

1. Registration Number

2. Name or corporate name of registrant

3. Permanent or legal address of registrant

4. Address(es) of site(s) where registered activities take place

5. National legal basis of registration

6. Name of responsible officer of the competent authority of the member state validating the registration

7. Date

Regulation 327 of The Human Medicines Regulations 2012 (SI 2012/1916)

EUROFINS SELCIA LIMITED, FYFIELD BUSINESS AND RESEARCH PARK, FYFIELD ROAD, ONGAR, CM5 0GS, UNITED KINGDOM

EUROFINS SELCIA LIMITED, FYFIELD BUSINESS AND RESEARCH

PARK, FYFIELD ROAD, ONGAR, CM5 0GS, UNITED KINGDOM

Confidential

13/05/2024

This registration form is valid only when presented with all pages. The authenticity of this registration form may be verified in MHRA-GMDP.

The registration holder referred to in section 2 shall communicate annually to the competent authority an inventory of the changes which have taken place as regards the information provided in this registration form. Any changes that may have an impact on the quality or safety of the listed active substances must be notified immediately.

SCOPE OF REGISTRATION

Name and address of the site

EUROFINS SELCIA LIMITED, FYFIELD BUSINESS AND RESEARCH PARK, FYFIELD ROAD, ONGAR, CM5 0GS, UNITED KINGDOM

1. MANUFACTURING OPERATIONS

Active substance

ACTIVE SUBSTANCES FOR CLINICAL TRIALS

А	Manufacture of Active Substance by Chemical Synthesis		
	A.1 Manufacture of Active Substance Intermediates		
	A.2 Manufacture of Crude Active Substance		

	A.3 Salt Formation / Purification Steps (e.g. Crystallisation) Preparative LC, crystallisation,	
E	General Finishing Steps	
	E.2 Primary Packaging	
	E.3 Secondary Packaging	
F	Quality Control Testing	
	F.1 Physical / Chemical testing	
	RA	JRF