



Medicines & Healthcare products
Regulatory Agency

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Regulatory Agency**

10 South Colonnade
Canary Wharf
London
E14 4PU
United Kingdom

gov.uk/mhra

Mayfair Laboratories Limited
23 Stanton Avenue
Manchester, England
M20 2PG
England, United Kingdom

16 August 2025

Dear **Gokhan Karahan**

We are writing to inform you of the outcome of the application to register or update an existing registration for the following manufacturer, which you submitted on **16 August 2025**:

Application reference: **2025081601433700**

Manufacturer organisation: **Mayfair Laboratories Limited**

Address:

23 Stanton Avenue
Manchester, England
M20 2PG
England, United Kingdom

Manufacturer registration status: **Registered**

Device(s):

GMDN Code & Term	Status	Comment
61087 - Clinical management support software	Registered	

Important Information:

Where new devices or device amendments within this application have been accepted and the status indicates 'Registered', this email confirmation does not represent any form of accreditation, certification or approval by the UK Competent Authority.

Where new devices within this application have been rejected, please review the reason/s for rejection. You will need to submit a new application. The [statutory fee](#) will be payable. Placing medical devices on the GB market that have not been registered with MHRA, or no longer comply with the regulations, constitutes a breach of the law. Different regulations apply to [Northern Ireland](#).

The name and address of the manufacturer and UK Responsible Person or Northern Ireland Authorised Representative (where applicable) and devices that have been registered will be published on our [Public Access Registration Database](#) (PARDB). In vitro diagnostic medical devices registered as undergoing performance evaluation study are not published on this database.

Keeping your Device Registration record up to date:

Please see [Making changes to your registration](#) for further information on changes you need to notify MHRA of, and the applicable [statutory fee](#). Full instructions on how to make changes are explained in our [Reference Guides](#) and [video tutorials](#).

Note:

The account number for your company/organisation is **0000035855**. Please keep a record of this.

Please do not reply directly to this email, as the originating email account is not monitored. Any queries must be sent to device.registrations@mhra.gov.uk.

Yours sincerely,



Ngozi Onyeukwu
MHRA Device Registration Service
Data Assurance & Quality
Healthcare, Quality & Access Group
Medicines and Healthcare products Regulatory Agency
10 South Colonnade, Canary Wharf, London, E14 4PU
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