



Medicines & Healthcare products Regulatory Agency

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Viamed Limited 15 Station Road Cross Hills West Yorkshire Keighley BD20 7DT England, United Kingdom

29 December 2021

Dear Stephen Nixon

We are pleased to confirm that the application to register or update an existing registration for the following manufacturer, which you submitted on 29 December 2021 has been reviewed:

Application reference: 2021122901230478

Manufacturer organisation: Ontecklong (Nanjing) Medical Supply Corporation Limited

Address:
Building 5,
No. 2 Future Road
Park Zhongshan
Jiangsu
Nanjing
Liuhe District
China

Manufacturer registration status: Registered

Device(s):

GMDN term	Status	Comment
11661 - Eye pad	Registered	

Please note this letter does not represent any form of <u>accreditation</u>, <u>certification or approval</u> by the UK Competent Authority.

If you stop placing devices on the market or if you are not complying with the Regulations, you should inform us so that we can amend our records. You should be aware that it is an offence to place on the market UKCA or CE marked devices that do not comply with the regulations.

Please inform us of the following chargeable changes:

- 1. company/organisation information e.g. name and address
- 2. additional devices (GMDN code or term)

Please also use the Devices Online Registration Database (DORS) to tell us of the following changes e.g. removal/discontinuation of a device (GMDN) or product from your registration record, change of contact person, telephone number and/or email address, for which payment of our statutory fee does not apply.

Please note that the name and address of manufacturer, UK Responsible Person or Authorised Representative (Northern Ireland only) and devices that have been registered will be published on our <u>Public Access Registration</u> Database (PARD).

The account number for your company/organisation is **0000021624.**

Yours sincerely,

Ngozi Onyeukwu

Device registrations service

Myellen.

Devices division

MHRA