



Certificate of Compliance

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We hereby declare that the technical files of all the items in each product group complies with the requirements of the Council Directive on Medical Devices 93/42/EEC as Amended 2007/47/EC Class 1.

Certificate No: - CE-3226

Manufacturer: MEDOX BIOTECH INDIA PRIVATE LIMITED

Address: 6/3, KODAMBAKKAM ROAD WEST, WEST MAMBALAM,

CHENNAI, TAMIL NADU-600 033, INDIA

Products: 1) MOLECULAR BIOLOGY KITS & REAGENTS,

ANALYTICAL/TEACHING KITS, ENZYMES & BIO CHEMICALS.
2) LIQUID HANDLING, PLASTIC WARES, ELECTROPHORESIS,

POWER PACK, UV TRANSILLUMINATORS, GEL

DOCUMENTATION SYSTEMS, LIVE GEL VISUALIZATION

UNITS & OTHER BIOTECH ALLIED APPARATUS

Complies with the requirements applicable to it

The quality system file has been assessed, approved and is subject to continuous surveillance according to the Council Directive on Medical Devices 93/42/EEC as Amended 2007/47/EC Class 1.

This certificate is issued under the following conditions:

- 1. It applies only to the quality system maintained in the manufacture of above referenced models and it does not substitute the design or type-examination procedures, if requested.
- The certificate remains valid until the manufacturing conditions or the quality systems are changed.
- 3. The certificate validity is conditioned by positive results or surveillance audits.

The CE mark as shown above can be used, under the responsibility of the manufacturer, after completion of an EC Declaration of conformity and compliance with all relevant EC Directives. The statement is based on a single evaluation of test report of one sample of above mentioned product. It does not imply an assessment of the whole production.

Validity of this certificate can be verified at www.ukcertifications.org.uk/verify

Date of Certification20th June 20221st Surveillance Audit Due19th June 20232nd Surveillance Audit Due19th June 2024Certificate Expiry (subject to the company maintaining19th June 2025

its system to the required standard)



