



CERTIFICATE

EC Certificate No. 1434-IVDD-394/2020
Full Quality Assurance System

Directive 98/79/EC concerning
***in vitro* diagnostic medical devices**

Polish Centre for Testing and Certification certifies
that the quality assurance system in the organization:

Premier Medical Corporation Private Limited
A1-302, GIDC, Sarigam 396155, Dist. Valsad,
Gujarat, INDIA

for the design, manufacture and final inspection of *in vitro* diagnostic medical device
List A

First Response® HBsAg Card Test
Ref: PI10FRC05CE, PI10FRC10CE, PI10FRC25CE, PI10FRC30CE

complies with requirements
of Annex IV (excluding Section 4, 6) to Directive 98/79/EC (as amended)
implemented into Polish law,
as evidenced by the audit conducted by the PCBC

Validity of the Certificate: from 26.10.2020 to 27.05.2024

The date of issue of the Certificate: 26.10.2020



Issued under the Contract No. **MD-121/2019**
Application No: **185/2017**
Certificate bears the qualified signature.
Warsaw, 26.10.2020
Module **H7**

Anna
Małgorzata
Wyroba
Vice-President

Elektronicznie
podpisany przez Anna
Małgorzata Wyroba
Data: 2020.10.26
10:59:54 +01'00'



CERTIFICATE

EC Certificate No. 1434-IVDD-396/2020
Full Quality Assurance System

Directive 98/79/EC concerning
***in vitro* diagnostic medical devices**

Polish Centre for Testing and Certification certifies
that the quality assurance system in the organization:

Premier Medical Corporation Private Limited
A1-302, GIDC, Sarigam 396155, Dist. Valsad,
Gujarat, INDIA

for the design, manufacture and final inspection of *in vitro* diagnostic medical device
List A

First Response® HCV Card Test
Ref: PI03FRC05CE, PI03FRC10CE, PI03FRC25CE, PI03FRC30CE

complies with requirements
of Annex IV (excluding Section 4, 6) to Directive 98/79/EC (as amended)
implemented into Polish law,
as evidenced by the audit conducted by the PCBC

Validity of the Certificate: from 26.10.2020 to 27.05.2024

The date of issue of the Certificate: 26.10.2020



Issued under the Contract No. **MD-121/2019**
Application No: **186/2017**
Certificate bears the qualified signature.
Warsaw, 29.09.2020
Module **H7**

Anna
Małgorzata
Wyroba
Elektronicznie
podpisany przez Anna
Małgorzata Wyroba
Data: 2020.10.26
11:10:39 +01'00'
Vice-President



सत्यमेव जयते

FORM MD-9

[See sub-rule (1) rule 25]

Licence to Manufacture for Sale or for Distribution of Class C or Class D medical device

Licence Number: MFG/MD/2018/000064

Endorsement No. 1

1. M/s PREMIER MEDICAL CORPORATION PRIVATE LIMITED, A1-302 GIDC Sarigam Dist, Valsad, Gujarat (India) - 396155 Telephone No.: 260 2780112, 260 2780113 FAX: 260 2242411 has been licenced to manufacture for sale or for distribution the below listed medical device(s) at the premises situated at M/s PREMIER MEDICAL CORPORATION PRIVATE LIMITED, A1-302, GIDC, Sarigam, Valsad, Gujarat (India) - 396155 Telephone No.: 260-2780112, 260-2780113 FAX: 260-2242411
2. Details of medical device(s) [Annexed]
3. The names, qualifications and experience of the competent technical staff responsible for the manufacture and testing of the above mentioned medical device(s): As per records maintain by the manufacturer
4. This licence is subject to the provisions of the Medical Devices Rules, 2017 and conditions prescribed therein.

ANNEXURE

| S.No. | Details Of Device(s) |
|-------|----------------------|
|-------|----------------------|

सत्यमेव जयते

**S ESWARA
REDDY**

Digitally signed by S ESWARA REDDY
DN: c=IN, o=CENTRAL DRUGS STANDARD
CONTROL ORGANIZATION,
2.5.4.20=cc9e6dd3ecf4ca36d3ae9e432d59a
6f4c3a02cb467e76d5edcb3a731ee144,
ou=DGHS,CID=6432120, postalCode=110002,
st=Delhi, cn=S ESWARA REDDY
Date: 2019.07.15 16:53:04 +05'30'

Certificate of Registration

QUALITY MANAGEMENT SYSTEM - ISO 13485:2016 & EN ISO 13485:2016

This is to certify that:

Premier Medical Corporation Private
Limited
A1-302, GIDC
Sarigam
Dist. Valsad 396 155
Gujarat
India

Holds Certificate No:

MD 584949

and operates a Quality Management System which complies with the requirements of ISO 13485:2016 & EN ISO 13485:2016 for the following scope:

The Design, Development and Manufacture of In-vitro Diagnostics Kits for Diagnosis of Infectious Diseases.

For and on behalf of BSI:

Gary E Slack, Senior Vice President - Medical Devices

Original Registration Date: 2006-02-20

Latest Revision Date: 2021-08-03

Effective Date: 2021-02-21

Expiry Date: 2024-02-20

Page: 1 of 1



...making excellence a habit.™

This certificate was issued electronically and remains the property of BSI and is bound by the conditions of contract.

An electronic certificate can be authenticated [online](#).

Printed copies can be validated at www.bsi-global.com/ClientDirectory or telephone +91 11 2692 9000.

Further clarifications regarding the scope of this certificate and the applicability of ISO 13485:2016 & EN ISO 13485:2016 requirements may be obtained by consulting the organization.

This certificate is valid only if provided original copies are in complete set.



Tel. direct: +41 22 791 3927
Fax direct: +41 22 791 4836
E-mail : diagnostics@who.int

In reply please
refer to: CC/vl

Your reference: P17-370-9

Premier Medical Corporation Private Limited
Attention: Dr Rajeshkumar Patel
Department of General Management
1304 Johnston Drive
Watchung, New Jersey
07069
Etats Unis-d' Amerique

24 June 2019

Dear Dr Patel,

Subject: WHO Prequalification of In Vitro Diagnostics – Final Public Report

Product name: First Response® HIV 1+2/Syphilis Combo Card Test
Product codes: I20FRC25, I20FRC30, I20FRC50, I20FRC60 and I20FRC100
Regulatory version: Rest of World
Manufacturer: Premier Medical Corporation Private Limited
PQDx Reference Number: PQDx 0364-010-00

We are pleased to inform you that the above-referenced product was prequalified on 24 June 2019 and listed on the World Health Organization (WHO) list of prequalified in vitro diagnostic products.

The following post-prequalification activities are required to maintain the prequalification status:

1. Notification to WHO of any planned changes to a prequalified product, in accordance with “WHO procedure for changes to a WHO prequalified in vitro diagnostic” (document number PQDx_121); and
2. Post-market surveillance activities, in accordance with “WHO guidance on post-market surveillance of in vitro diagnostics” (ISBN 978 92 4 150921 3).

ENCL: as stated

You are also required to submit an annual report that details sales data and all categories of complaints in a summarized form. There are certain categories of complaints and changes to the product that must be notified immediately to WHO, as per the above-mentioned documents. The sales data will serve as denominator data to guide the frequency of re-inspection.

Failure to comply with any of the above-mentioned post-prequalification requirements may lead to remedial action by WHO, including but not limited to, de-listing from the WHO list of prequalified in vitro diagnostic products.

If you have any questions, please do not hesitate to contact us by email (diagnostics@who.int) or by telephone (+4122 791 3927).

Yours sincerely,



Mr Deus Mubangizi
Coordinator
Prequalification Team
Regulation of Medicines and other Health Technologies



Tel. direct: +41 22 791 3927
Fax direct: +41 22 791 4836
E-mail : diagnostics@who.int

In reply please
refer to: CC/vl

Your reference: P17-370-9

Premier Medical Corporation
Private Limited
Attention: Dr Rajeshkumar Sheliya
Director MR
A1-302, GIDC
Sarigam, District, Valsad
396155
Inde

13 January 2021

Dear Dr Sheliya,

Subject: WHO Prequalification of In Vitro Diagnostics – Final Public Report

Product name: First Response Syphilis Anti-TP Card Test

Product codes: PI08FRC25, PI08FRC50 and PI08FRC100

Regulatory version: Rest of World

Manufacturer: Premier Medical Corporation Private Limited

PQDx Reference Number: PQDx 0471-010-00

We are pleased to inform you that the above-referenced product was prequalified on 13 January 2021 and listed on the World Health Organization (WHO) list of prequalified in vitro diagnostic products.

The following post-prequalification activities are required to maintain the prequalification status:

1. Notification to WHO of any planned changes to a prequalified product, in accordance with “WHO procedure for changes to a WHO prequalified in vitro diagnostic” (document number PQDx_121); and
2. Post-market surveillance activities, in accordance with “WHO guidance on post-market surveillance of in vitro diagnostics” (ISBN 978 92 4 150921 3).

You are also required to submit an annual report that details sales data and all categories of complaints in a summarized form. There are certain categories of complaints and changes to the product that must be notified immediately to WHO, as per the above-mentioned documents. The sales data will serve as denominator data to guide the frequency of re-inspection.

Failure to comply with any of the above-mentioned post-prequalification requirements may lead to remedial action by WHO, including but not limited to, de-listing from the WHO list of prequalified in vitro diagnostic products.

If you have any questions, please do not hesitate to contact us by email (diagnostics@who.int) or by telephone (+4122 791 3927).

Yours sincerely,



Mr Deus Mubangizi
Unit Head
Prequalification Unit
Regulation and Prequalification Department

WHO list of prequalified in vitro diagnostic products

RoW: Rest of the world. Regulatory version applied to products not approved by stringent/mature NRAs or not regulated
 Last update: 22 February 2022

| Year prequalified | Type of assay | Product name | Product code(s) | Regulatory version | Manufacturer | Manufacturing site(s) | Packaging |
|-------------------|--------------------------|--|--|--------------------|---|---|---|
| 2019 | HBSAg RDT | *Determine HBSAg 2 | 7D2942; 7D2943; 7D2943 SET | CE-mark | Alere Medical Co. Ltd | 357 Matsuhidai, Matsudo-shi, 270-2214, Chiba-ken, Japan | 20 T/kit 100 T/kit 100 T/kit |
| 2019 | HCV EIA | ARCHITECT HCV Ag assay | 6L47-29; 6L47-11; 6L47-02; and 8C89-01 | CE-mark | Denka Seiken Co., LTD, Kagamida Factory | Street 1359-1, Kagamida, Kigoshi, Gosen-shi, Niigata, Japan | 100 T/kit |
| 2019 | HIV RDT for self-testing | *Mylan HIV Self Test | ARST001-03; ARST001-03-01; ARST001-03-02; ARST001-03-03 | RoW | Atomo Diagnostics Pvt. Ltd | Site 1: Atomo Diagnostics Pty Ltd at Level 2, 701-703 Parramatta Road, Leichardt 2040 NSW, Australia Site 2: Lateral Flow Laboratories (LFL) at Unit 1 & 2, Greenwich Place, Capricorn Crescent, Capricorn Technology Park, Muizenberg, 7945, South Africa | 1 T/kit; 1 T/kit; 1 T/kit; 1 T/kit. |
| 2019 | HIV/Syp RDT | *First Response HIV1+2/Syphilis Combo Card Test | I20FRC25; I20FRC30; I20FRC50; I20FRC60; I20FRC100 | RoW | Premier Medical Corporation Private Limited | Sarigam, Gujarat, India | 25 T/kit 30 T/kit 50 T/kit 60 T/kit 100 T/kit |
| 2019 | HIV RDT | *ONE STEP Anti-HIV (1&2) Test | ITPW02152-TC40; ITPW02152-TC25; ITPW02153-TC40 ITPW02153-TC40SA | RoW | InTec PRODUCTS, INC | 308, Wengjiao Rd, Xinyang IND. AREA, Haicang, Xiamen, 361022, China | 40 T/kit 25 T/kit 40 T/kit 40 T/kit |
| 2019 | HCV RDT | Rapid Anti-HCV Test | ITPW01152-TC40; ITPW01152-TC25; ITPW01153-TC40 | RoW | InTec PRODUCTS, INC | 308, Wengjiao Rd, Xinyang IND. AREA, Haicang, Xiamen, 361022, China | 40 T/kit 25 T/kit 40 T/kit |
| 2019 | Malaria RDT | AdvDx Malaria Pf Rapid Malaria Ag Detection Test | 00-DKM-RK-MALADX-004-025 | RoW | Advy Chemical Pvt Ltd., | Plot No.A-334,336,338 & A-337 & 339 Road no. 25 & 26, Wagle industrial Estate Thane 400 604 India | 25 T/kit |
| 2019 | Malaria RDT | *NxTek Eliminate Malaria Pf | 05FK140 | CE-mark | Abbott Diagnostics Korea Inc | site 1: 46, Hagal-ro 15 beongil, Giheung-gu, Yongin-si, Gyeonggi-do 17099, Republic of Korea site 2: 65, Borahagal-ro, Giheung-gu, Yongin-si, Gyeonggi-do 17099, Republic of Korea | 25T/kit |
| | | | 05FK141 | | | | 25T/kit |
| | | | 05FK142 | | | | 1T/kit x 25 each |
| | | | 05FK143 | | | | 1T/kit x 25 each |
| 2019 | HIV NAT | *m-PIMA HIV-1/2 VL | 27015-W50 | RoW | Abbott Rapid Diagnostics Jena GmbH | Orlaweg 1, D-07743 Jena, Germany | 50 cartridges/kit |
| | | First Response Malaria Antigen P | PI13FRC25s | | | site 1: A1-302, GIDC, Sarigam 396 155, Valsad, Gujarat, India; | 25 × single kit |
| | | | PI13FRC10s | | | | 10 × single kit |