



Food and Drug Administration
10903 New Hampshire Avenue
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Silver Spring, MD 20993-0002

August 12, 2014

Remote Diagnostic Technologies Ltd.
Leigh Cornock
Director Of Systems Engineering & Test
The Old Coach House, The Avenue
Farleigh Wallop, Basingstoke
Hampshire, RG25 2HT UK

Re: K134014
Trade/Device Name: Tempus Pro Accessories (ultrasound & video laryngoscope probe)
Regulation Number: 21 CFR 870.2300
Regulation Name: Cardiac Monitor (Including Cardiotachometer and Rate Alarm)
Regulatory Class: Class II
Product Code: MWI, IYO, ITX
Dated: June 25, 2014
Received: July 3, 2014

Dear Dr Leigh Cornock,

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the [Federal Register](#).

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); medical device reporting (reporting of medical device-

related adverse events) (21 CFR 803); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please contact the Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address

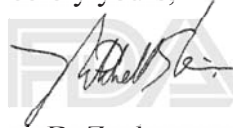
<http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm>. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to

<http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm> for the CDRH's Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

You may obtain other general information on your responsibilities under the Act from the Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address

<http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm>.

Sincerely yours,

A handwritten signature in black ink, appearing to read "Bram D. Zuckerman", is written over a large, light gray watermark of the FDA logo.

for Bram D. Zuckerman, M.D.

Director

Division of Cardiovascular Devices

Office of Device Evaluation

Center for Devices and Radiological Health

Enclosure

Indications for Use Statement

510(k) Number (if known): K134014

Device Name: Tempus Pro

Indications for Use:

The Tempus Pro is a portable vital signs monitor intended to be used by clinicians and medically qualified personnel for the attended or unattended monitoring of single or multiple vital signs in clinical and pre-hospital care applications. The device is indicated for 3 & 5 Lead ECG monitoring and 12 Lead ECG recording, impedance pneumography, non-invasive blood pressure (NIBP), end-tidal CO₂ (ETCO₂) and respiration rate, pulse oximetry (SpO₂), contact temperature and invasive pressure.

The monitor is intended to be used as a stand-alone monitor or as a telemedicine system (transmitting patient data to other medical professionals located elsewhere).

The device is indicated for adults, paediatrics and neonates.

The monitor can be used to display images from Interson 3.5 MHz General Purpose (GP) and 7.5 MHz Small Parts / Vascular (SR) USB ultrasound probes or a Karl Storz C-MAC S USB video laryngoscope. These optional accessories are to be used in accordance with their Indications For Use.


Prescription Use _____
(Part 21 CFR 801 Subpart D)

AND/OR

Over-The-Counter Use _____
(21 CFR 801 Subpart C)

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of Device Evaluation (ODE)

 Date:
2014.08.12
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510(k) Summary of Safety and Effectiveness

5.1 Submitter's Information

The submitter of this special pre-market notification is:

Name: Remote Diagnostic Technologies Limited
 Address: The Old Coach House, Farleigh Wallop, Basingstoke,
 RG25 2HT, United Kingdom
 Company Phone No: +44 (0) 1256 362 400
 Company Fax No: +44 (0) 1256 362 415
 Contact Person: Dr Leigh Cornock (Director of Systems Eng. & Test)
 Date summary prepared: 23rd December, 2013

5.2 Device Identification

Device Trade Name: Tempus Pro
 Common Name: Patient Monitor (without arrhythmia detection or alarms)
 Class: II
 Classification Panel: 74
 Product Code: MWI
 Secondary Product Codes: IYO & ITX
 Regulation Number: 870.2300

5.3 Device Description

The Tempus Pro is a multi-parameter vital signs monitor designed for use in pre-hospital care and remote clinical locations by trained healthcare professionals. It provides 3&5 lead ECG monitoring, 12 lead ECG recording, pulse oximetry, non-invasive blood pressure, sidestream capnometry, contact temperature, impedance respiration, invasive pressure and user configurable alarms.

In addition, it provides the ability to transmit all vital signs data via wired or wireless connections to a software system (called i2i) expected to be based in a facility far from the user e.g. a response centre facility. In addition to sending all vital signs, the system can also send pictures or video via an integrated camera, geographic position by an integrated GPS receiver and voice via a wired or wireless headset.

5.4 Indications for Use

The following indications for use of the existing features of Tempus Pro Patient Monitor remain unchanged by addition of the option to used plug-in probes for displaying ultrasound and video laryngoscopy images on the monitor.

"The Tempus Pro is a portable vital signs monitor intended to be used by clinicians and medically qualified personnel for the attended or unattended monitoring of single or multiple vital signs in clinical and pre-hospital care applications. The device is indicated for 3 & 5 Lead ECG monitoring and 12 Lead ECG recording, impedance pneumography, non-invasive blood pressure (NIBP), end-tidal CO₂ (ETCO₂) and respiration rate, pulse oximetry (SpO₂), contact temperature and invasive pressure.

The monitor is intended to be used as a stand-alone monitor or as a telemedicine system (transmitting patient data to other medical professionals located elsewhere).

The device is indicated for adults, paediatrics and neonates.

The monitor can be used to display images from Interson 3.5 MHz General Purpose (GP) and 7.5 MHz Small Parts / Vascular (SR) USB ultrasound probes or a Karl Storz C-MAC S USB video laryngoscope. These optional accessories are to be used in accordance with their Indications For Use."

Remote Diagnostic Technologies Ltd - Tempus Pro™ Patient Monitor with Accessories 510(k)

5.5 Comparison with Cleared Device

The intended use and indications for use, plus the fundamental technology used in the Tempus Pro device, remain essentially unchanged as a result of plugging in the optional accessories, the monitor may be used to display ultrasound and video laryngoscopy images.

5.6 Substantial Equivalence

The predicate for the Tempus Pro Patient Monitor with the specified accessories attached is the cleared Tempus Pro Patient Monitor (K130773).

The Predicate for the optional Ultrasound features are:

- K070907 – Interson USB ultrasound probe system
- K102153 – MobiUS Ultrasound imaging system

As the accessories are unmodified, claims for substantial equivalence relate to the display of images, using drivers supplied by the OEM (where appropriate) and additional RDT software to enable their display on the existing Tempus Pro screen.

5.8 Summary of Non-Clinical Testing

Addition of optional probes connected externally to Tempus Pro and changes to the associated software have been made under well-established design control procedures, which ensure that appropriate risk management processes have been carried out to determine their impact and ensure that appropriate verification/ validation testing is performed.

The non-clinical testing carried out in relation to addition of the optional imaging probes described above is summarized in the following table:

Area	Testing Performed
Comparative Testing to predicates	Comparative testing has been performed to demonstrate that the performance of the device is equivalent to the predicates.
Software	The requirements of the FDA document <i>Guidance for the Content of Premarket Submissions for Software in Pre-Market Submissions</i> has been applied. In addition, the requirements of IEC 62304 have been addressed.
Bench testing	All parameters of the device have been tested to confirm they operate to specification across their stated performance range and across their stated temperature range with the new accessories attached.
Bench testing	The product has been bench tested to confirm that all data is transmitted reliably and accurately.

In each case the results of this testing confirmed that acceptance criteria defined by the relevant standard, or other appropriate reference document had been met.

With respect to usability, no additional user validation was considered necessary, as the Tempus Pro used with these accessories is identical to the predicate Tempus Pro in terms of physical and interface features, including size and weight, user interface data layouts, button styles, menus, and layout of connectors, resulting the physical interaction with the device being unchanged. The way the accessories are used is un-changed as a result of using them with the Tempus Pro. .

5.8 Conclusion

In accordance with the Federal Food, Drug and Cosmetic Act, 21 CFR Part 807 and based on the information provided in this Premarket Notification, Remote Diagnostic Technologies Limited conclude that the Tempus Pro Patient Monitor when used with optional probes for ultrasound and video laryngoscopy is safe and effective, and substantially equivalent to the unmodified version of this device.