--- ELECTRICAL BUSINESS GROUP - ELECTRONIC PRODUCTS -

Mysore Works, KIADB Industrial Area, Hebbal - Hootagalli, Mysore - 570 018 • Tel : (91) - 821 - 2402561 • Fax : (91) - 821 - 2402468

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Ref.

JUN 2 9 2005

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510(K) SUMMARY (Per section 807.92 ©)

CONTACT DAT	ГА		
Submitter's Name Address		Larsen & Toubro Limited KIADB Industrial Area, Hebbal Hootagalli, Mysore - 570018, Karnataka, INDIA	
Contact Person	A.B.Deshpande	Title	Head - Quality Assurance
E-Mail address		DeshpandeAB@myw.ltindia.com	
Date the summary was prepared		April 25 th ,2004	

Regd. Off : L & T House, Ballard Estate, P. O. Box 278, Mumbai 400 001 • Phone | 261 8181 / 82 • Fax · 91 - 22 - 262 0223 Website: www.intebg.com



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DEVICE				
Trade name	STAR 50			
Соттоп пате	Patient Monitoring System			
Classification name	Vital Signs Monitor			

PREDICATE DEVICE IDENTIFICATION						
CFR21 Section 870.2300	Product code (optional) MWI					
Classification panel	Cardiovascular					
Device Class	Class II					
Legally marketed Comparison Device / K#	Eagle 3000 patient Monitoring System (Marquette Electronic) / K952474					
	 Vital signs monitor Model 8100 (CSI) / K001020 					
	 Patient Monitoring System - STAR, (Larsen & Toubro Limited), K# K032867 					



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DEVICE DESCRIPTION

This STAR 50 unit is a multiparameter Patient monitor System (TFT color monitor) with ECG(3/5 lead), Respiration, Temperature, NIBP, Pulse oximetry, Capnography and Invasive BP.

STAR 50 is a four channel monitor with waveform display capability for ECG (Lead I / III / V / AVL / AVF / AVR), Plethysmograph, Respiration, Invasive Blood pressure (IBP1 & IBP2) and Capnography (CO₂). It also displays the digital values of HR/PR, SpO₂, RR, Non-Invasive & Invasive Blood Pressure (Systolic, Diastolic and Mean), Temperature, EtCO₂ and FiCO₂ readings. It has graded and color coded alarms. It has 24 hours tabular and graphical trends for all parameters except NIBP. For NIBP the last 240 readings tabular trend can be seen. Display of last 16 alarm conditions is possible in alarm recall mode. STAR 50 has got a optional Thermal recorder for printing Tabular trends & waveforms.

INTENDED USE OF THE DEVICE

The STAR 50 multiparameter Patient Monitoring system is intended to monitor a single adult, pediatric or neonatal patient's vital signs at the bedside or during intra-hospital transport along with the appropriate accessories mentioned / supplied with the unit. Vital signs parameters include ECG (Lead I / II / III / V / AVL / AVF / AVR), Plethysmograph, Respiration, Invasive Blood pressure (IBP1 & IBP2) and Capnography (CO₂). It can also display the digital values of HR/PR, SpO₂, RR, Non-Invasive & Invasive Blood Pressure (Systolic, Diastolic and Mean), Temperature, EtCO₂ and FiCO₂ readings.

The user, responsible to interpret the monitored data made available, will be a professional health care provider. The device, which can also be used as a portable device, permits patient monitoring with adjustable alarm limits as well as visible and audible alarm signals. The monitor is not intended for home use.

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TECHNOLOGICAL CHARACTERISTICS OF THE DEVICE COMPARED TO THE PREDICATE DEVICE

Device: Larsen & Toubro limited make STAR 50 Patient Monitoring System.

Predicate device:

Eagle 3000 patient Monitoring System (Marquette Electronic), K# K952474 Vital signs monitor Model 8100 (CSI), K# K001020 Patient Monitoring System – STAR, (Larsen & Toubro Limited), K# K032867

The parameters available with these predicate devices are available with the Larsen & Toubro Limited make STAR 50 patient monitoring system (ECG-3/5 lead, Respiration, Temperature – 2 channels, NIBP, Pulse oximetry, Capnography and Invasive BP- 2 channels). The no. of channels, range and accuracy of the parameters & method of sensing are similar to the predicate devices. In STAR 50 monitor audible & visual alarms are provided similar to those in the Predicate devices.

STAR 50 has got TFT color display like CSI Model 8100. Weight is also comparable with that of Marquette Eagle 3000. Battery (2 scaled lead acid) is provided in STAR 50 monitor like that of the predicate device CSI Model 8100.

Comparison of all the parameters of STAR 50 to that of the predicate devices is given in the "Substantial Equivalence Equipment comparison" document.

Compliance to standards:

The following international standards are referred.

IEC 60601-1 Medical Electrical safety

IEC 60601-1-2 EMC compliance

IEC 60601-2-27 ECG safety

Conclusion:

Based on the Technological characteristics of STAR50 and its comparison with those of a predicate device CSI Model 8100 and Marquettee Eagle 3000 monitors, Larsen & Toubro Limited believes that their device is substantially equivalent to these Monitors and doesn't pose any additional tisk on safety & effectiveness of the device.

(N Ravindran)

Head - Design & Development

Regd. Off: L. & T. House, Ballard Estate, P. O. Box 278, Mumbai 400.001 • Phone: 261.8181 / 82 • Fax: 91 - 22 - 262.0223. Website: www.Intebg.com



Food and Drug Administration 9200 Corporate Boulevard Rockville MD 20850

JUN 2 9 2005

Larsen & Toubro Ltd. c/o Mr. Ned Devine Intertek Testing Services NA, Inc. 3033 Madison Ave. SE Grand Rapids, MI 49548

Re: K051608

Trade Name: STAR 50 Monitoring System Regulation Number: 21 CFR 870.2300

Regulation Name: Physiological Patient Monitor without Arrhythmia Detection and Alarms

Regulatory Class: Class II (two)

Product Code: MWI Dated: June 16, 2005 Received: June 17, 2005

Dear Mr. Devine:

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.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to such 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.

Page 2 -Mr. Ned Devine

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); 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. This letter will allow you to begin marketing your device as described in your Section 510(k) premarket notification. The FDA finding of substantial equivalence of your device to a legally marketed predicate device results in a classification for your device and thus, permits your device to proceed to the market.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please contact the Office of Compliance at (240) 276-0295 Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97) you may obtain. Other general information on your responsibilities under the Act may be obtained from the Division of Small Manufacturers, International and Consumer Assistance at its toll-free number (800) 638-2041 or (301) 443-6597 or at its Internet address http://www.fda.gov/cdrh/dsma/dsmamain.html

Sincerely yours,

Bram D. Zuckerman, M.D.

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Director

Division of Cardiovascular Devices

Office of Device Evaluation

Center for Devices and

Radiological Health

Enclosure

Indications for Use

510(k) Number (if known): <u>K051608</u>

Device Name: STAR 50

Indications for Use:

The STAR 50 unit is a multi-parameter Patient Monitoring system is intended to monitor a single adult, pediatric or neonatal patient's vital signs at the bedside or during intra-hospital transport along with the appropriate accessories mentioned/supplied with the unit. Vital signs parameters include ECG (Lead I / III / V / AVL / AVF AVR), Plethysmograph, Respiration, Invasive Blood Pressure (IBP1 & IBP2) and Capnography (CO₂). It can also display the digital values of HR/PR, SpO₂, RR, Non-Invasive & Invasive Blood Pressure (systolic, diastolic and mean) and Temperature, EtCO₂ and FiCO₂ readings.

The user, responsible to interpret the monitored data made available, will be a professional health care provider. The device, which can also be used as a portable device, permits patient monitoring with adjustable alarm limits as well as visible and audible alarm signals. The arrhythmia provided is only a rate related arrhythmia without alarms. The monitor is not intended for home use.

Prescription Use		A
(Part 21 CFR 801	Subpart D)	А

AND/OR

Over-The-Counter Use _____ (21 CFR 807 Subpart C)

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

Concurrence of CDRH, Office of Device Evaluation (ODE)

(Division Sign-Off)

Division of Cardiovascular Devices

510(k) Number KOS/6AX