

CE DECLARATION OF CONFORMITY

LeadCare® Plus™ BloodLead System

Manufacturer: Hersteller, Fabricante Fabricant, Produttore	Fabricante, Producent Tillverkare, Κατασκευαστής	Magellan Diagnostics 101 Billerica Ave, Building 4 North Billerica, Massachusetts 01862 U.S.A
EU Authorized Representative: EU-Bevollmächtigte Representante Autorizado por la UE Mandataire Rappresentante Autorizzato in Eu	Representante Autorizado na UE EU- autoriseret repræsentant EU Auktoriserad representant Εξουσιοδοτημένος αντιπρόσωπος στην ΕΕ	Ichor Technologies Ltd 1 Paper Mews, 330 High Street Dorking, Surrey, RH4 2TU, UK

Magellan Diagnostics hereby declares that the products listed below conform to the European Union directive and standards identified in this declaration.

Magellan Diagnostics erkl art, dass die aufgef hrten Produkt(e) mit den Bestimmungen der angegebenen EU-Richtlinien und mit den aufgef hrten normativen Dokumenten in  bereinstimmung sind.

Magellan Diagnostics declara por la presente que los producto(s) abajo mencionados, est n conformes con las directivas y normas Europeas identificadas en esta declaraci n.

Magellan Diagnostics d clare par la pr sente, que le(s) produit(s) sous-mentionn (s), est (sont) conforme(s) aux directives et normes Europ ennes identifi es dans cette d claration.

Magellan Diagnostics dichiara con la presente che il(i) prodotto(i) sottomenzionato(i)  (sono) conformi alla direttiva e agli standard specificati in questa dichiarazione.

Magellan Diagnostics declara pelo presente que o(s) produto(s) abaixo mencionado(s) est /est o conforme a Directiva e normas da Comiss o Europeia especificadas nesta declara o.

Magellan Diagnostics erkl rer herved, at det (de) nedenfor anf rte produkt(er) er i overensstemmelse med de EU-direktiver og standarder, der er anf rt i denne erkl ring.

Magellan Diagnostics bekr ftar h rmed att produkt(er) listade nedan, vara f renlig(a) med Europeiska Union-ens direktiv och standarder identifierade i denna deklaration.

H Magellan Diagnostics με το παρόν δηλώνει ότι τα προϊόντα που αναφέρονται κατωτέρω συμμορφούνται με την οδηγία της Ευρωπαϊκής Ένωσης και τα πρότυπα που προσδιορίζονται στην παρούσα δήλωση.

EU Directive:

EU-Richtlinie Directiva UE Directive Europ enne Direttiva Europea Directiva UE EU-direktiv EU Direktiv Οδηγία ΕΕ

IVD - 98/79/EC (27/10/1998) – Annex I and III, Excluding Section 6 for Self Test

- ISO 9001:2008, Quality Management System
- ISO 13485:2003, Medical Devices – Quality Systems
- 21CFR Part 809.10, 812.5, 820.130-820.160(a), 820.30, 820.70
- ISO 14971:2007, Medical Devices - Application of risk management to medical devices
- EN 61010-1:2001 (2cd Edition),(IEC 1010-1) : Safety requirements for electrical equipment for measurement, control, and laboratory use - Part 1: General requirements
- EN 61010-2-101 Part 2: Particular Requirements for in-vitro diagnostic (IVD) medical equipment.
- EN 61326:2002, Group 1, Class B , Electrical equipment for measurement, control and laboratory use – EMC requirements, Amendment 1:1998 – Immunity ; Industrial
- ISO 15223:2007, Medical Devices – Symbols to be used with medical devices, labelling and information to be supplied
- EN 980:2008, Graphical symbols for use in the labeling of medical devices
- EN 375:2001, Information supplied by the manufacturer with in vitro diagnostic reagents for professional use
- EN 18113-3:2011, Instructions for use for in vitro diagnostic instruments for professional use
- EN ISO 23640:2011 In vitro diagnostic medical devices; Evaluation of stability of in vitro diagnostic reagents


 Director of QA/RA


 Date

Product(s)		Beginning	
Produkt(e)	Produto(s)	zu beginnen von	Inicio
Producto(s)	Produkt(er)	A partir de	Gældende fra
Produit(s)	Produkt(er)	Première id.	From
Prodotto(i)	Προϊόντα	A partire da	Εναρξη

Product	P/N	GMDN CODE	GMDN CODE	
LeadCare® Plus™ Analyzer System	82-0001	31646	21 batt01 10 01	July 2015
LeadCare® Plus™ Blood Lead Test Kit	82-0004	31646	21 batt01 10 01	