



Declaration of Conformity

PRODUCT IDENTIFICATION CHRONIC INTESTINAL DISEASE FAMILY OF PRODUCTS	
Product name	Model/number
IBD-CHEK®	T5008, 30352
IBD-SCAN®	T5009, 30351
ASCA-CHEK	T5016, 30361
IBD EZ VUE®	30353
LEUKO EZ VUE®	T30355
Kits listed include all of the corresponding components.	

MANUFACTURER		
Name of company	Address	Representative
TECHLAB®	2001 Kraft Dr Blacksburg, VA 24060 U.S.A.	Donna T. Link – Director Regulatory and Compliance and Quality Management Representative

AUTHORIZED REPRESENTATIVE		
Name of company	Address	Telephone/email
Emergo Europe	Molenstraat 15 2513 BH The Hague, Netherlands	+31.70.345.8570 - phone +31.70.346.7299 - fax service@emergogroup.com

CONFORMITY ASSESSMENT		
Device classification	Route to compliance	Standards applied
Self-Certify	Annex III of IVDD 98/79/EEC Council Directive	ISO 13485:2003

TECHLAB® declares that the above mentioned products meet the provision of the Council Directive 98/79/EEC for *in vitro* diagnostic Medical Devices.

COMPANY REPRESENTATIVE:
TITLE:

Donna T. Link
Director Regulatory and Compliance, QMR

SIGNATURE:

DATE:

2014/08/22
YYYY/MM/DD