



CE Technical Documentation Review Report

Applicant: Sheyang Huida Medical Products Co., Ltd.
Qinfeng Industrial Park, Yandong, Sheyang, Jiangsu
224300, China

Report Number: 15018206 001

Examination intent: Examination the completeness of the Technical Documentation according to the requirements of the In Vitro Diagnostic Medical Devices Directive 98/79/EC Annex III

Product(s): Microscope Slides, Microscope Cover Slips

Type(s)/Model(s): Various

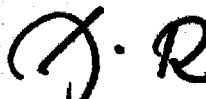
Classification: IVD other product
(according to manufacturer's declaration)

Examination period: 2006 July 17

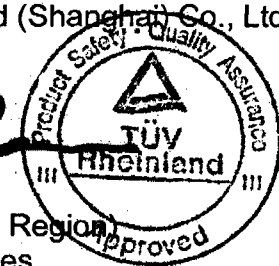
Date of expiry: 16.07.2011

Review result: During the examination of the provided Technical Documentation (HD/CE-001, Revision: A/0, Date 2006-06-15) no Non-compliance according to the requirements of the In Vitro Diagnostic Medical Devices Directive 98/79/EC Annex III was detected.

TÜV Rheinland (Shanghai) Co., Ltd.


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Manager (Asia Region)
Medical Services



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