
	EU Declaration of Conformity accordingly Directive 98/79/EG, Annex III	Revision: 14/01.02.2017 Dok.-Nr.: 30705/ EN-UA
		1 / 1

Manufacturer: Ahlstrom Germany GmbH
Niederschlag 1
09471 Bärenstein
Deutschland
T: +49 (0) 37347 83 - 0
F: +49 (0) 37347 83 - 64

Product name: TFN-Specimen Collection Card

Item group: 2.460.00054

Indented use: Specimen collection paper for absorption, storage and transport of human sample blood (dry blood spot sample) for In-Vitro-diagnostics of Newborn screenings and HIV1 analysis.



Classification: General IVD
EDMS 26-02 (sample processor)

Declaration: Within the meaning of Annex III of Directive 98/79/EC we declare as manufacturer with sole responsibility that the specimen collection paper mentioned above fulfills all essential requirements of Directive 98/79/EC of the European Parliament and the Council of 27.10.1998 on In vitro diagnostics.

Harmonised standard: DIN EN ISO 13485:2012-11
DIN EN 980:2008-08
DIN EN 13612:2002-08
DIN EN ISO 14971:2013-04
DIN EN ISO 18113-1:2013-01

Product standard: The requirements of the product standard CLSI document NBS01-A6 - Blood Collection on Filter Paper for Newborn Screening Programs are fulfilled.

Validity: This document is valid until 31.01.2018.

		
Date Signature	11.05.2017	11.05.2017
Name Function	Thomas Foltyn Plant Manager	Maria-Isabel Buhler Quality Manager

In the case of unauthorized modifications in the product or a not indented use this declaration becomes invalid.