

info@fdab.com e-mail www.fdab.com website SE556450496601 vat no. Göteborg place o

e-mail website vat no. place of registered office Fujirebio Diagnostics AB Majnabbeterminalen SE-414 55 Göteborg, Sweden +46 31 85 70 30 phone +46 31 85 70 40 fax

CERTIFICATE OF CONFORMITY

We, Fujirebio Diagnostics AB hereby declare that the products listed below comply with the In Vitro Medical Device Directive 98/79/EC and its relevant transposition into the national laws of the member states in which the devices are intended to be placed on the market.

	Prod no	Prod grouping
CanAg CA242 EIA	101-10*	Common/Other IVD product
CanAg CA19-9 EIA	120-10*	Common/Other IVD product
CanAg CA15-3 EIA	200-10*	Common/Other IVD product
CanAg PSA EIA	340-10**	Annex II list B
CanAg Free PSA EIA	350-10**	Annex II list B
CanAg CA125 EIA	400-10*	Common/Other IVD product
CanAg CEA EIA	401-10*	Common/Other IVD product
CanAg NSE EIA	420-10*	Common/Other IVD product
CanAg AFP EIA	600-10*	Common/Other IVD product
CanAg S100 EIA	708-10*	Common/Other IVD product
CanAg SCC EIA	800-10*	Common/Other IVD product
CanChek Tumor Marker Control Serum	107-20*	Common/Other IVD product
HE4 EIA	404-10*	Common/Other IVD product
Cyfra 21-1 EIA	211-10*	Common/Other IVD product

* using Annex III as the conformance assessment procedure

** using Annex IV as the conformance assessment procedure.

December 1, 2009 Göteborg

Fujirebio Diagnostics AB

all

Christina Hall Managing Director



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tel fax e-mail website

DECLARATION OF CONFORMITY

We, Fujirebio Diagnostics AB hereby declare that the products listed below comply with the In Vitro Medical Device Directive 98/79/EC and its relevant transposition into the national laws of the member states in which the devices are intended to be placed on the market.

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CanAg CA19-9 EIA	120-10*	Common/Other IVD product
CanAg CA242 EIA	101-10*	Common/Other IVD product
CanAg CEA EIA	401-10*	Common/Other IVD product
CanAg Free PSA EIA	350-10**	Annex II list B
CanAg NSE EIA	420-10*	Common/Other IVD product
CanAg ProGRP EIA	220-10*	Common/Other IVD product
CanAg PSA EIA	340-10**	Annex II list B
CanAg S100 EIA	708-10*	Common/Other IVD product
CanAg SCC EIA	800-10*	Common/Other IVD product
CanChek	107-20*	Common/Other IVD product
CYFRA 21-1 EIA	211-10*	Common/Other IVD product
HE4 EIA	404-10*	Common/Other IVD product
Tumor Marker Control	108-20**	Annex II list B
ProGRP Control	230-20*	Common/Other IVD product
Mesothelin Control	360-20*	Common/Other IVD product
Lung Marker Control	240-20*	Common/Other IVD product

* using Annex III as the conformance assessment procedure

** using Annex IV as the conformance assessment procedure.

16 February 2016, Göteborg

Fujirebio Diagnostics AB

Thomas Si

Thomas Stjernkvist OA & RA Manager

EC Certificate TÜVRheinland Directive 98/79/EC Annex IV, excluding Sections 4 and 6 Full Quality Assurance System In Vitro Diagnostic Medical Devices				
	Registration No.:	HL 60139396	6 0001	
	Report No.:	21220990 00	09	FWIREBIO
Manufacturer:	Fujirebio Diagnostics Elof Lindälvs gata 1 SE-414 58 Göteborg Sweden	3	om e	Thomas Stjernikvist, M. Sc. Regulatory Affairs & Quality Systems Manager Fujirebio Diagnostics AB Elof Lindälvs gata 13 SE-414 58 Göteborg, Sweden Phone: + 46 (0) 31 304 90 00 www.fdab.com - info@fdab.com
Products:	Reagents and reagent the tumoral marker E	-	termining	
	Replaces Certificate	e, Registration N	O.: HL 60096	5837 0001
Expiry Date:	2024-05-24			
The Notified Body hereby declares that the requirements of Annex IV, excluding section 4 and 6 of the directive 98/79/EC have been met for the listed products. The above named manufacturer has established and applies a quality assurance system, which is subject to periodic surveillance, defined by Annex IV, section 5 of the aforementioned directive. For placing on the market of List A devices covered by this certificate an EC design-examination certificate according to Annex IV, section 4 and 4 LGA Andrew verification of manufactured products according to section 6 is required.				
Effective Date:	2019-10-16		\leq	UVRheinland
Date:	2019-07-18		DiplIng. Sten	Hoffmann
TÜV Rheinland LGA Products GmbH - Tillystraße 2 - 90431 Nürnberg TÜV Rheinland LGA Products GmbH is a Notified Body according to Directive 98/79/EC concerning in vitro diagnostic medical devices with the identification number 0197.				



Fujirebio Diagnostics AB Elof Lindälvs gata 13 Box 121 32 SE-414 58 Göteborg Sweden

To whom it concern

FREE SALES CERTIFICATE

It is hereby certified that Fujirebio Diagnostics AB, Elof Lindälvs gata 13, Box 121 32, SE-414 58 Göteborg, Sweden, is the manufacturer of the product/products listed in the attached product list.

The Swedish Medical Products Agency certifies that these devices are not under restrictions given by the Swedish Medical Product Agency and may be exported without restrictions. The devices are freely marketed in Sweden.

This certificate is valid until May 25, 2022

On behalf of the Swedish Medical Products Agency



Postadress/Postal address: P.O. Box 26, SE-751 03 Uppsala, SWEDEN Besöksadress/Visiting address: Dag Hammarskjölds väg 42, Uppsala Telefon/Phone: +46 (0)18 17 46 00 Fax: +46 (0)18 54 85 66 Internet: www.lakemedelsverket.se E-mail: registrator@lakemedelsverket.se



Product Name	Article Number
AMH Control	660-20 / 17350066140579
CanAg AFP EIA	600-10 / 17350066140371
CanAg CA125 EIA	400-10 / 17350066140241
CanAg CA15-3 EIA	200-10 / 17350066140081
CanAg CA19-9 EIA	120-10 / 17350066140050
CanAg CA242 EIA	101-10 / 17350066140012
CanAg CEA EIA	401-10 / 17350066140289
CanAg Free PSA EIA	350-10 / 17350066140203
CanAg NSE EIA	420-10 / 17350066140340
CanAg ProGRP EIA	220-10 / 17350066140159
CanAg PSA EIA	340-10 / 17350066140173
CanAg S100 EIA	708-10 / 17350066140388
CanAg SCC EIA	800-10 / 17350066140401
CanChek Tumor Marker Control	107-20 / 17350066140036
CYFRA 21-1 EIA	211-10 / 17350066140135
HE4 EIA	404-10 / 17350066140302
Lung Marker Control	240-20 / 17350066140531
Mesothelin Control	360-20 / 17350066140524
Tumor Marker Control	108-20 / 17350066140043



Fujirebio Diagnostics AB PO Box 121 32 SE-402 42 Göteborg Sweden

delivery/visiting address Elof Lindälvs gata 13 SE-414 58 Göteborg Sweden

vat no. SE556450496601 place of registered office Göteborg

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DECLARATION OF CONFORMITY

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Lung Marker Control	240-20*	Common/Other IVD product
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* using Annex III as the conformance assessment procedure

** using Annex IV as the conformance assessment procedure.

22 January 2021, Göteborg

Fujirebio Diagnostics AB

Juan Sig

Thomas Stjernkvist Regulatory affairs and Quality systems Manager



Thomas Stjernkvist, M. Sc. Regulatory Affairs & Quality Systems Manager

Fuji rebio Diagnostics AB Elof Lindālvs gata 13 SE-414 58 Göteborg, Sweden

Phone: + 48 (0) 31 304 90 00 www.tdab.com - info@fdab.com



CERTIFICATE

The Certification Body of TÜV SÜD Management Service GmbH

certifies that



Fujirebio Germany GmbH Hans-Böckler-Allee 20 30173 Hannover Germany

has established and applies a Quality Management System for

Sales, customer services and marketing of in vitro diagnostics and instruments.

An audit was performed, Order No. 707118450.

Proof has been furnished that the requirements according to

ISO 9001:2015

are fulfilled. The certificate is valid from 2020-06-19 until 2023-06-18. Certificate Registration No.: 12 100 59925 TMS.

Product Compliance Management Munich, 2020-05-06



CERTIFICATE

ERTIFIKAT 🔶







Certificate No. Q5 083208 0032 Rev. 03

Holder of Certificate:

Fujirebio Inc.

2-1-1 Nishishinjuku Shinjuku-ku, Tokyo 163-0410 JAPAN

Certification Mark:



Scope of Certificate:

Design and Development, Production and Distribution of Immunoassay Reagent Kits, Reagent Kits for Biochemistry, Substrate Solution, Wash Solution, Specimen Diluent and Control Design and Development, Production and Distribution of Instruments for Immunoassay, Sampling Tips and Specimen Container Installation and Servicing of Instruments for Immunoassay

The Certification Body of TÜV SÜD Product Service GmbH certifies that the company mentioned above has established and is maintaining a quality management system, which meets the requirements of the listed standard(s). All applicable requirements of the testing and certification regulation of TÜV SÜD Group have to be complied with. For details and certificate validity see: www.tuvsud.com/ps-cert?q=cert:Q5 083208 0032 Rev. 03

Report No.:

JN1667218

Valid from: Valid until: 2021-09-06 2024-09-05

Date,

2021-09-02

Christoph Dicks Head of Certification/Notified Body





Certificate

No. Q5 083208 0032 Rev. 03

Applied Standard(s): EN ISO 13485:2016 Medical devices - Quality management systems -Requirements for regulatory purposes (ISO 13485:2016) DIN EN ISO 13485:2016

Facility(ies):Fujirebio Inc.2-1-1 Nishishinjuku, Shinjuku-ku, Tokyo, 163-0410 JAPAN

Design and Development, Production and Distribution of Immunoassay Reagent Kits, Reagent Kits for Biochemistry, Substrate Solution, Wash Solution, Specimen Diluent and Control Distribution of Instruments for Immunoassay, Sampling Tips and Specimen Container

Fujirebio Inc. Hachioji Facility 51, Komiya-machi, Hachioji-shi, Tokyo, 192-0031 JAPAN

Design and Development, Production and Distribution of Immunoassay Reagent Kits, Reagent Kits for Biochemistry, Substrate Solution, Wash Solution, Specimen Diluent and Control Design and Development, Production and Distribution of Instruments for Immunoassay, Sampling Tips and Specimen Container Installation and Servicing of Instruments for Immunoassay

Fujirebio Inc. Hachioji 2nd Facility 153, Komiya-machi, Hachioji-shi, Tokyo, 192-0031 JAPAN

Production and Distribution of Immunoassay Reagent Kits, Substrate Solution, Wash Solution, Specimen Diluent and Control

Fujirebio Inc. Hachioji 3rd Facility 935, Ishikawa-machi, Hachioji-shi, Tokyo, 192-0032 JAPAN

Production and Distribution of Immunoassay Reagent Kits, Reagent Kits for Biochemistry, Substrate Solution, Wash Solution, Specimen Diluent and Control

Fujirebio Inc. Sagamihara Facility 1-3-14, Tanashioda, Chuo-ku, Sagamihara-shi, Kanagawa, 252-0245 JAPAN

Production and Distribution of Immunoassay Reagent Kits, Substrate Solution, Wash Solution, Specimen Diluent and Control





Certificate No. Q5 083208 0032 Rev. 03

Facility(ies):

Fujirebio Inc. Tokachi Obihiro Facility 8-36, Nishi-3-sen, Aza-Otofuke, Otofuke-cho, Katou-gun, Hokkaido, 080-0341 JAPAN

Production and Distribution of Immunoassay Reagent Kits, Substrate Solution, Wash Solution, Specimen Diluent and Control

Fujirebio Inc. Ube Facility 203-152, Aza-Ushiake, Oaza-Yoshiwa, Ube-shi, Yamaguchi, 759-0134 JAPAN

Production and Distribution of Immunoassay Reagent Kits, Substrate Solution, Wash Solution, Specimen Diluent and Control

Fujirebio Inc. Distribution Center 1-17-19, Haijima-cho, Akishima-shi, Tokyo, 196-0002 JAPAN

Distribution of Immunoassay Reagent Kits, Reagent Kits for Biochemistry, Substrate Solution, Wash Solution, Specimen Diluent and Control Distribution of Instruments for Immunoassay, Sampling Tips and Specimen Container

Fujirebio Inc. Kuki Distribution Center 1662, Shimohayami, Kuki-shi, Saitama, 346-0022 JAPAN

Distribution of Immunoassay Reagent Kits, Reagent Kits for Biochemistry, Substrate Solution, Wash Solution, Specimen Diluent and Control Distribution of Instruments for Immunoassay, Sampling Tips and Specimen Container

Fujirebio Inc. Asahikawa Facility 23-1975-167, Minamiyonjodori, Asahikawashi, Hokkaido, 078-8334 JAPAN

Production and Distribution of Immunoassay Reagent Kits, Substrate Solution, Wash Solution, Specimen Diluent and Control

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