

MANUFACTURER	DIA.PRO DIAGNOSTIC BIOPROBES S.R.L.
	VIA G. CARDUCCI Nº 27 – 20099 SESTO SAN
	GIOVANNI (MILANO) – ITALY
PRODUCT	CMV IgG
x	CODE: CMVG.CE (96 tests)
CLASSIFICATION	ANNEX II – LIST B
CONFORMITY ASSESSMENT ROUTE	ANNEX IV

#### WE HEREBY DECLARE THAT THE ABOVE MENTIONED PRODUCT MEETS THE PROVISIONS OF THE COUNCIL DIRECTIVE 98/79/EC FOR IN VITRO DIAGNOSTIC DEVICES.

NOTIFIED BODY	$AEMPS - n^{\circ} 0318$
(EC) CERTIFICATE(S)	<ul> <li>FULL QUALITY ASSURANCE SYSTEM N°</li> </ul>
	2004 05 0442 CT (in accordance with Annex IV -
	except Section IV) of the Directive 98/79/EC),
	RELEASED BY EC NOTIFIED BODY N° 0318
	• UNI EN ISO 13485 N° 2013 11 0039 EN,
	RELEASED BY EC NOTIFIED BODY N° 0318

PLACE & DATE OF FIRST ISSUE	MILANO – MAY 2004
PLACE & DATE OF CURRENT	SESTO SAN GIOVANNI (MI) – DECEMBER 2013
EMISSION	
SIGNATURE	
Legal Representative	DIA. PRO. DIAGNOSTIC BLOPBOBES SI
Dr.ssa Fiorenza Scozzesi	DIAGROSTIC DIOPRODES SI
	Addee
0	

Rev: 12/2013



MANUFACTURER	DIA.PRO DIAGNOSTIC BIOPROBES S.R.L.
	VIA G. CARDUCCI Nº 27 – 20099 SESTO SAN
	GIOVANNI (MILANO) – ITALY
PRODUCT	CMV IgM
	CODE: CMVM.CE (96 tests)
CLASSIFICATION	ANNEX II – LIST B
<b>CONFORMITY ASSESSMENT ROUTE</b>	ANNEX IV

#### WE HEREBY DECLARE THAT THE ABOVE MENTIONED PRODUCT MEETS THE PROVISIONS OF THE COUNCIL DIRECTIVE 98/79/EC FOR IN VITRO DIAGNOSTIC DEVICES.

NOTIFIED BODY	AEMPS – n° 0318
(EC) CERTIFICATE(S)	<ul> <li>FULL QUALITY ASSURANCE SYSTEM N°</li> </ul>
	2004 05 0442 CT (in accordance with Annex IV –
	except Section IV) of the Directive 98/79/EC),
	RELEASED BY EC NOTIFIED BODY N° 0318
	• UNI EN ISO 13485 N° 2013 11 0039 EN,
	RELEASED BY EC NOTIFIED BODY N° 0318

PLACE & DATE OF FIRST ISSUE	MILANO – MAY 2004
PLACE & DATE OF CURRENT	SESTO SAN GIOVANNI (MI) – DECEMBER 2013
EMISSION	16 152
SIGNATURE	
Legal Representative	DIA. PRO. DIAGNOSTIC BIOPROBES STI
Dr.ssa Fiorenza Scozzesi	DIAGNOSTIC DIOPROBES SI
	Lotte
	H

Rev: 12/2013

DIA.PRO Diagnostic Bioprobes S.r.I. Sede legale e lab.: Via G.Carducci, 27 – 20099 Sesto S.Giovanni (MI) – Italia Tel. +39 02 27007161/6450 • Fax +39 02 26007726 • http://www.diapro.it • E-mail: info@diapro.it Capitale sociale €50.000,00 I.V. – P.IVA: 11924660159 – Reg. Imp. 11924660159 – REA 1509959



DIPARTIMENTO DELLA PROGRAMMAZIONE E DELL'ORDINAMENTO DEL SERVIZIO SANITARIO NAZIONALE DIREZIONE GENERALE DEI DISPOSITIVI MEDICI, DEL SERVIZIO FARMACEUTICO E DELLA SICUREZZA DELLE CURE UFFICIO IV ex DGFDM – DIAGNOSTICI IN VITRO

#### I.5.I.e.2/IV/2011/37

**VISTA** la direttiva 98/79/CE relativa ai dispositivi medico-diagnostici in vitro; **VISTO** il D.lgs. n .332/2000 recante attuazione della direttiva 98/79/CE;

**VISTA** l'istanza del 29/09/2011 presentata dalla ditta Dia.Pro Diagnostic Bioprobes Srl con sede in Via G.Carducci, 27 – 20099 Sesto San Giovanni (MI) – C.F./P.Iva

11924660159; CONSIDERATO che la ditta istante ha effettuato i versamenti richiesti dal D.M. 24 Maggio 2004;

**VISTI** gli atti d'ufficio;

**HAVING REGARD** to 98/79/EC directive concerning the in vitro diagnostic medicaldevices;

**HAVING REGARD** to legislative Decree (D.lgs.)n. 332/2000 reporting the accomplishment of 98/79/EC Directive;

**HAVING REGARD** to *the request dated 29/09/2011 submitted by the company* Dia.Pro Diagnostic Bioprobes Srl con with legal site in Via Columella, 31 – 20128 Milano – C.F. and P.Iva 11924660159;

**WHEREAS** this company paid the fees required by Ministerial Decree (D.M.) May 24, 2004; **HAVING REGARD** to the official deeds;

#### SI ATTESTA IT IS ATTESTED

che la ditta, Dia. Pro Diagnostic Bioprobes Srl con sede in Via G.Carducci, 27 – 20099 Sesto San Giovanni (MI) – C.F./P.Iva 11924660159, ha prodotto e marcato CE, come dispositivo medico- diagnostico in vitro, secondo le procedure previste dalla direttiva 98/79/CE, il prodotto:

that the Company Dia.Pro Diagnostic Bioprobes Srl located in Via G.Carducci, 27 - 20099Sesto San Giovanni (MI) – C.F./P.Iva 11924660159, manufactured and affixed CE marking as in vitro diagnostic medical device, according to the Directive 98/79/EC, the following product:

#### DP-9 DIA.BLOOD INSTRUMENT

Il suddetto prodotto, in base all'art. 4 della direttiva 98/79/CE, è di libera circolazione e può essere messo in commercio in Italia e in tutto il territorio dell'Unione Europea.



Si rilascia il presente attestato su richiesta dell'interessato per gli usi consentiti dalla legge e per l'esportazione nei paesi extra UE.

The above mentioned product, according to the art. 4 of 98/79/EC directive, can freely circulate and can be commercialized in Italy and in the whole of the European Union. This certificate is issued on the interested company's request according to the law and to export to non-European countries

LASALUF IL DIRETTORE DELL'UFFICIO IV (Dott.ssa Giovanna Nisticò) varino Instic RTIMEN

IC/CM



MANUFACTURER	DIA.PRO DIAGNOSTIC BIOPROBES S.R.L.
	VIA G. CARDUCCI Nº 27 – 20099 SESTO SAN
	GIOVANNI (MILANO) – ITALY
PRODUCT	TOXO IgG
	CODE: TOXOG.CE (96 tests)
CLASSIFICATION	ANNEX II – LIST B
<b>CONFORMITY ASSESSMENT ROUTE</b>	ANNEX IV

#### WE HEREBY DECLARE THAT THE ABOVE MENTIONED PRODUCT MEETS THE PROVISIONS OF THE COUNCIL DIRECTIVE 98/79/EC FOR IN VITRO DIAGNOSTIC DEVICES.

NOTIFIED BODY	$AEMPS - n^{\circ} 0318$
(EC) CERTIFICATE(S)	• FULL QUALITY ASSURANCE SYSTEM N° 2004 05 0442 CT (in accordance with Annex IV –
	except Section IV) of the Directive 98/79/EC), RELEASED BY EC NOTIFIED BODY N° 0318
	<ul> <li>UNI EN ISO 13485 N° 2013 11 0039 EN, RELEASED BY EC NOTIFIED BODY N° 0318</li> </ul>

PLACE & DATE OF FIRST ISSUE	MILANO – MAY 2004
PLACE & DATE OF CURRENT	SESTO SAN GIOVANNI (MI) – DECEMBER 2013
EMISSION	
SIGNATURE	
Legal Representative	DIA_PRO
Dr.ssa Fiorenza Scozzesi	DIA PRO. DIAGNOSTIC BIOPROBES STI
	Hydro
	100 lot L

Rev: 12/2013



MANUFACTURER	DIA.PRO DIAGNOSTIC BIOPROBES S.R.L.
	VIA G. CARDUCCI Nº 27 – 20099 SESTO SAN
	GIOVANNI (MILANO) – ITALY
PRODUCT	TOXO IgM
	CODE: TOXOM.CE (96 tests)
CLASSIFICATION	ANNEX II – LIST B
<b>CONFORMITY ASSESSMENT ROUTE</b>	ANNEX IV

#### WE HEREBY DECLARE THAT THE ABOVE MENTIONED PRODUCT MEETS THE PROVISIONS OF THE COUNCIL DIRECTIVE 98/79/EC FOR IN VITRO DIAGNOSTIC DEVICES.

NOTIFIED BODY	$AEMPS - n^{\circ} 0318$
(EC) CERTIFICATE(S)	<ul> <li>FULL QUALITY ASSURANCE SYSTEM N° 2004 05 0442 CT (in accordance with Annex IV – except Section IV) of the Directive 98/79/EC), RELEASED BY EC NOTIFIED BODY N° 0318</li> <li>UNI EN ISO 13485 N° 2013 11 0039 EN, RELEASED BY EC NOTIFIED BODY N° 0318</li> </ul>

PLACE & DATE OF FIRST ISSUE	MILANO – MAY 2004
PLACE & DATE OF CURRENT	SESTO SAN GIOVANNI (MI) – DECEMBER 2013
EMISSION	
SIGNATURE	DIA. PRO. DIAGNOSTIC BIOPROBES STI
Legal Representative Dr.ssa Fiorenza Scozzesi	DIAGNOSTIC BIOPROBES STI
	Aforthe (

Rev: 12/2013



Lorne Laboratories Ltd declares that the following in vitro diagnostic reagent:

Product name	Catalogue number
ASO Latex kit	031100A

has been classified as non List A, non List B (Directive 98/79/EC, Annex II) and complies with the essential requirements and provisions of Directive 98/79/EC of the European Parliament and of the Council (also SI 2002 No.618 which transposes the requirements of Directive 98/79/EC).

and is in conformity with the national standards transposing harmonised standards:

- BS EN 980:2008
- BS EN ISO 13485:2012
- BS EN 13612:2002
- BS EN 13640:2002
- BS EN 13641:2002
- BS EN ISO 14971:2012
- BS EN ISO 18113, parts 1&2

The conformity assessment procedure performed was in accordance with Annex III of Directive 98/79/EC.

This declaration of conformity is issued under the sole responsibility of Lorne Laboratories Ltd and is valid from 13 April 2016.

S

Eddy Velthuis Technical Director



Lorne Laboratories Limited Unit 1 Cutbush Park Industrial Estate Danehill, Lower Earley Berkshire RG6 4UT United Kingdom

Tel: +44 (0) 118 921 2264 Fax: +44 (0) 118 986 4518 Email: info@lornelabs.com www.lornelabs.com



Lorne Laboratories Ltd declares that the following in vitro diagnostic reagent:

Product name	Catalogue number
RF Latex kit	830100A

has been classified as non List A, non List B (Directive 98/79/EC, Annex II) and complies with the essential requirements and provisions of Directive 98/79/EC of the European Parliament and of the Council (also SI 2002 No.618 which transposes the requirements of Directive 98/79/EC).

and is in conformity with the national standards transposing harmonised standards:

- BS EN 980:2008
- BS EN ISO 13485:2012
- BS EN 13612:2002
- BS EN 13640:2002
- BS EN 13641:2002
- BS EN ISO 14971:2012
- BS EN ISO 18113, parts 1&2

The conformity assessment procedure performed was in accordance with Annex III of Directive 98/79/EC.

This declaration of conformity is issued under the sole responsibility of Lorne Laboratories Ltd and is valid from 13 April 2016.

S

Eddy Velthuis Technical Director



Lorne Laboratories Limited Unit 1 Cutbush Park Industrial Estate Danehill, Lower Earley Berkshire RG6 4UT United Kingdom

Tel: +44 (0) 118 921 2264 Fax: +44 (0) 118 986 4518 Email: info@lornelabs.com www.lornelabs.com



Lorne Laboratories Ltd declares that the following in vitro diagnostic reagent:

Product name	Catalogue number
CRP Latex kit	850100A

has been classified as non List A, non List B (Directive 98/79/EC, Annex II) and complies with the essential requirements and provisions of Directive 98/79/EC of the European Parliament and of the Council (also SI 2002 No.618 which transposes the requirements of Directive 98/79/EC).

and is in conformity with the national standards transposing harmonised standards:

- BS EN 980:2008
- BS EN ISO 13485:2012
- BS EN 13612:2002
- BS EN 13640:2002
- BS EN 13641:2002
- BS EN ISO 14971:2012
- BS EN ISO 18113, parts 1&2

The conformity assessment procedure performed was in accordance with Annex III of Directive 98/79/EC.

This declaration of conformity is issued under the sole responsibility of Lorne Laboratories Ltd and is valid from 13 April 2016.

S

Eddy Velthuis Technical Director



Lorne Laboratories Limited Unit 1 Cutbush Park Industrial Estate Danehill, Lower Earley Berkshire RG6 4UT United Kingdom

Tel: +44 (0) 118 921 2264 Fax: +44 (0) 118 986 4518 Email: info@lornelabs.com www.lornelabs.com



## **EC CERTIFICATE**

### Lorne Laboratories Ltd

Unit 1 Cutbush Park Industrial Estate, Danehill, Lower Earley, Berkshire RG6 4UT, UK

### EC Certificate - Full Quality Assurance System Approval Certificate

Annex IV, (excluding sections 4 and 6) of Council Directive 98/79/EC on In Vitro Diagnostic Medical Devices

Scope of Certificate: The design and manufacture of in vitro diagnostic reagents for identification of blood groups

Device Classification: Annex II, List A and B

Device Descriptions: Please refer to Attachment 1

Model: Please refer to Attachment 1

File Number A12241 Certificate No. 354.170425 Cycle Start Date23 May 2017Effective Date23 May 2017Expiry Date22 May 2022

Authorised by

B. Rodgers Certification Manager For and on Behalf of UL International (UK) Ltd

We hereby declare that an examination of the full quality assurance system has been carried out per report 11640248 , following the requirements of the national legislation to which the undersigned is subject, transposing Annex IV (with the exemption of sections 4 and 6) of Council Directive 98/79/EC on In Vitro Diagnostic Medical Devices. We certify that the full quality assurance system conforms with the relevant provisions of the aforementioned directive and is subject to periodic surveillance as required by 98/79/EC, Annex IV, Section 5. For Annex II, List A devices where they are covered by this certificate, an EC Design Examination certificate according to 98/79/EC, Annex IV, Section 4 is required. This certificate is issued with 1 attachment listing model numbers.

Notified Body 0843

UL International (UK) Limited Wonersh House, The Guildway, Old Portsmouth Road, Guildford, Surrey, GU3 1LR, United Kingdom

## **EC CERTIFICATE**



### Lorne Laboratories Ltd

Unit 1 Cutbush Park Industrial Estate, Danehill, Lower Earley, Berkshire RG6 4UT, UK

#### Attachment 1 of 1

The products detailed below are covered under the scope of this certificate

Device Description	Model	Classification
Anti-A Monoclonal	600005/600010/600000	Annex II List A
Anti-B Monoclonal	610005/610010/610000	Annex II List A
Anti-A,B Monoclonal	620005/620010/620000	Annex II List A
Anti-C Monoclonal	690005	Annex II List A
Anti-E Monoclonal	691005	Annex II List A
Anti-c Monoclonal	692005	Annex II List A
Anti-e Monoclonal	693005	Annex II List A
Anti-K Monoclonal	760005/760010	Annex II List A
Anti-D Clone 2 Monoclonal	710010/710000	Annex II List A
Anti-D Clone 1 Monoclonal	730010/730000	Annex II List A
Anti-D Duoclone Monoclonal	740010/740000	Annex II List A
Anti-Jka Polyclonal	323002/323000	Annex II List B
Anti-Jkb Polyclonal	324002/324000	Annex II List B
Anti-Fyb Polyclonal	317002/317000	Annex II List B
AHG Elite Clear	415010/415100/415000	Annex II List B
AHG Elite Green	435010/435100/435000	Annex II List B
Anti-Fya Monoclonal	774000/774002	Annex II List B
Anti-C+D+E Monoclonal	700005/700010/700000	Annex II List A
Anti-Human IgG Clear	401010/401000	Annex II List B
Anti-Human IgG Green	402010/402000	Annex II List B
Monoclonal Rh Control	640010	Annex II List A
Monoclonal D Negative Control	650010	Annex II List A

File Number A12241 Certificate No. 354.170425 Cycle Start Date 23 May 2017 Effective Date 23 May 2017 Expiry Date 22 May 2022

Authorised by

B. Rodgers Certification Manager For and on Behalf of UL International (UK) Ltd

Notified Body 0843

IVDD A4 S3 FQ 00-NB-F0051 Issue: 6.0

UL International (UK) Limited Wonersh House, The Guildway, Old Portsmouth Road, Guildford, Surrey, GU3 1LR, United Kingdom