

Formularul ofertei (F3.1)

Data: 05.08-11.08.19

Nr.: 21010578

Alternativa Nr.: *nu sunt*

Către: AGENTIA NATIONALA pentru SANATATE PUBLICA

“GBG-MLD”SRL declară că:

- a) Au fost examinate și nu există rezervări față de documentele de atribuire.
- b) “GBG-MLD”SRL se angajează să furnizeze/presteze, în conformitate cu documentele de atribuire și condițiile stipulate în specificațiile tehnice și preț, următoarele bunuri și/sau servicii: - Denumirea-Achizitionarea truse pentru diagnosticul prin tehnici de biologie moleculara(PCR) si ELISA pentru anul 2019
- c) **Suma totală a ofertei fără TVA constituie: 49240,00**
[patruzeci și nouă mii doua sute patruzeci] Lei 00 bani.
- d) **Suma totală a ofertei cu TVA constituie: 58188.00**
[cincizeci și opt mii una suta optzeci și opt] Lei 00 bani.
- e) Prezenta ofertă va rămâne valabilă pentru perioada de timp specificată în **FDA4.8.**, începînd cu data-limită pentru depunerea ofertei, în conformitate cu **FDA5.2.**, va rămîne obligatorie și va putea fi acceptată în orice moment pînă la expirarea acestei perioade;
- f) În cazul acceptării prezentei oferte, “GBG-MLD”SRL se angajează să obțină o Garanție de bună execuție în conformitate cu **FDA7**, pentru executarea corespunzătoare a contractului de achiziție publică.
- g) Nu sîntem în nici un conflict de interese, în conformitate cu punctul **IPO5.4**.
- h) Compania semnatară, afiliații sau sucursalele sale, inclusiv fiecare partener sau subcontractor ce fac parte din contract, nu au fost declarate neeligibile în baza prevederilor legislației în vigoare sau a regulamentelor cu incidență în domeniul achizițiilor publice, în conformitate cu punctul **IPO5.5**.

Semnat: _____

L.Ș.

Nume: Tudor Ceaicovschi

În calitate de: director

Ofertantul: “GBG-MLD”SRL

Adresa: mun. Chisinau, str. Tighina 65 of. 607

Data: 05.08-11.08.19

Specificații tehnice (F4.1)

Numărul procedurii de achiziție: 21010578 din 05.08.2019 -11.08.19
 Denumirea procedurii de achiziție: valoare mică

Pagina 1 din 2

Denumirea bunurilor/serviciilor	Modelul articolului	Țara de origine	Produce-cătorul	Specificarea tehnică deplină solicitată de către autoritatea contractantă	Specificarea tehnică deplină propusă de către ofertant	Standarde de referință
Bunuri						
Lot 1						
IgM Measles Virus (Rujeola)	EI 2610-9601M	Germania	Euroimmun	vezi caietul de sarcini	Tip reacție - imunoenzimatică, pentru diagnostic uman. Durata perioadei de incubare în reacția de testare - până la 120 minute. Incubarea nu va include procesul de agitare. Lichid stabilii gata de lucru. Prezența în trusă a tuturor reactivelor necesare pentru reacție, inclusiv controalele. Placa de 96 godeuri (12 stripuri a câte 8 godeuri demonstrabile) Reactivi pentru 96 investigații inclusiv controalele. Sensibilitate testului nu mai puțin de 100 %, specificitate diagnostică nu mai puțin de 98 %.	CE,ISO
IgG Measles Virus (Rujeola)	EI 2610-9601G	Germania	Euroimmun	vezi caietul de sarcini	Tip reacție - imunoenzimatică, pentru diagnostic uman. Durata perioadei de incubare în reacția de testare - până la 120 minute. Incubarea nu va include procesul de agitare. Lichid stabilii gata de lucru. Prezența în trusă a tuturor reactivelor necesare pentru reacție, inclusiv controalele. Placa de 96 godeuri (12 stripuri a câte 8 godeuri demonstrabile) Reactivi pentru 96 investigații inclusiv controalele. Sensibilitate testului nu mai puțin de 100 %, specificitate diagnostică nu mai puțin de 100 %.	CE,ISO
Aviditatea IgG Measles virus (Rujeola)	EI 2610-9601-1G	Germania	Euroimmun	vezi caietul de sarcini	Tip reacție - imunoenzimatică. Principiul metodei - determinarea avidității anticorpilor către virusul rujeolei pentru diferențierea reinjecției de infecția primară. Lichid stabilii gata de lucru. Prezența în trusă a tuturor reactivelor necesare pentru reacție, inclusiv controalele și reagent pentru determinarea avidității. Sensibilitatea, și specificitatea $\geq 98\%$. Placa de 96 godeuri (12 stripuri a câte 8 godeuri demonstrabile). Reactivi pentru 96 investigații. CE pentru utilizare in vitro diagnostic.	CE,ISO
Lot 3						
Test ELISA Adenovirus Ag	1654Аденовирус-антиген-ИФА-БЕСТ,96 teste	Rusia	Vector Best	vezi caietul de sarcini	Lichid stabilii gata de lucru. Prezența în trusă a tuturor reactivelor necesare pentru reacție, inclusiv controalele. Placa de 96 godeuri (12 stripuri a câte 8 godeuri nedemonstrabile) Reactivi pentru 96 investigații inclusiv controalele.	ISO
Test ELISA Astrovirus Ag	EIA4456,Астровирус Аg (stool),96 teste	SUA	DRG	vezi caietul de sarcini	Lichid stabilii gata de lucru. Prezența în trusă a tuturor reactivelor necesare pentru reacție, inclusiv controalele. Placa de 96 godeuri (12 stripuri a câte 8 godeuri demonstrabile) Reactivi pentru 96 investigații inclusiv controalele.	ISO
Test ELISA Norovirus Ag	1656 Норовирус-антиген-ИФА-БЕСТ,96 teste	Rusia	Vector Best	vezi caietul de sarcini	Lichid stabilii gata de lucru. Prezența în trusă a tuturor reactivelor necesare pentru reacție, inclusiv controalele. Placa de 96 godeuri (12 stripuri a câte 8 godeuri nedemonstrabile) Reactivi pentru 96 investigații inclusiv controalele.	CE,ISO
Lot 4						
Coxiella burnetii Phase 1 IgG calitativ	COXI G0600, Coxiella Burnetii (Q-Fever) Phase 1 IgG,96 teste	Germania	Novatec	vezi caietul de sarcini	Lichid stabili gata de lucru. Prezența în trusă a tuturor reactivelor necesare pentru reacție, inclusiv controalele. Placa de 96 godeuri (12 stripuri a câte 8 godeuri demonstrabile) Reactivi pentru 96 investigații inclusiv controalele.	CE,ISO

Coxiella burnetii Phase 2 IgG cantitativ	COX2G0600 Coxiella Burnetii (Q-Fever) Phase 2 IgG	Germania	Novatec	vezi caietul de sarcini	Lichid stabilii gata de lucru. Prezența în trușă a tuturor reactivelor necesare pentru reacție, inclusiv controalele. Placa de 96 godeuri (12 stripurii a câte 8 godeuri demontabile) Reactivi pentru 96 investigații inclusiv controalele.	CE,ISO
Lot 34						
Test sistemă completă pentru determinarea ADN Coxiella burnetii în timp real (RT-PCR) în material biologic (singe, spună, spălături bronhice, lievor, material necrotic, etc.)	R-B85-50- F(R,G,i,Q,Mx,D) АмплиСенс® Coxiella burnetii-FL	Rusia	ЦНИИ Эпидемиол ологии	vezi caietul de sarcini	Reactivi pentru nu mai puțin de 50 teste, inclusiv controalele, comparabili cu amplificator Rotor-Gene-6000, (eprobete 0,2 ml). Sensibilitate între 1000-5000 copii/ml. Setul va include reactivi concepuți pentru a efectua o reacție PCR completă, care implică extracția ADN-ului din material biologic și amplificarea ADN-ului Coxiella burnetii cu detecție în timp real. Certificat CE pentru utilizarea in vitro diagnostic în conformitate cu cerințele Directivei Europene.	CE,ISO

Semnat: _____ Numele, prenumele: Ceaicovschi Tudor În calitate de: Director general

Ofertantul: „GBG-MLD” SRL Adresa: mun. Chișinău, str. Tighina, 65, of. 607

Specificații de preț (F4.2)

Numărul procedurii de achiziție: 21010578 din 05.08.2019 -11.08.19
 Denumirea procedurii de achiziție: valoare mica

Pagina 1 din 1

Cod CPV	Denumirea bunurilor/serviciilor	Unitatea de măsură	Cantitatea	Preț unitar (fără TVA)	Preț unitar (cu TVA)	Suma		Termenul de Livrare/prestare
						fără TVA	cu TVA	
33600000-6	Lot 1							
33600000-6	IgM Measles Virus (Rujeola)	trusă	1	3250,00	3900,00	3250,00	3900,00	30 zile de la solicitarea beneficiarului
33600000-6	IgG Measles Virus (Rujeola)	trusă	5	2593,00	3111,60	12965,00	15558,00	
33600000-6	Aviditatea IgG Measles virus (Rujeola)	trusă	2	3250,00	3900,00	6500,00	7800,00	
33600000-6	Total lot 1					6500,00	7800,00	
33600000-6	Lot 3							
33600000-6	Test ELISA Adenovirus Ag	trusă	1	1950,00	2340,00	1950,00	2340,00	30 zile de la solicitarea beneficiarului
33600000-6	Test ELISA Astrovirus Ag	trusă	1	7990,00	9588,00	7990,00	9588,00	
33600000-6	Test ELISA Norovirus Ag	trusă	1	2700,00	3240,00	2700,00	3240,00	
33600000-6	Total lot 3					2700,00	3240,00	
33600000-6	Lot 4							
33600000-6	Coxiella burnetii Phase 1 IgG calitativ	trusă	1	3750,00	4050,00	3750,00	4050,00	30 zile de la solicitarea beneficiarului
33600000-6	Coxiella burnetii Phase 2 IgG cantitativ	trusă	1	3750,00	4050,00	3750,00	4050,00	
33600000-6	Total lot 4					3750,00	4050,00	
33600000-6	Lot 7							
33600000-6	PCR test-sistemă completă pentru determinarea și diferențierea ADN Bordetella pertussis, Bordetella parapertussis și Bordetella bronchiseptica în regiun real time	set	1	6385,00	7662,00	6385,00	7662,00	30 zile de la solicitarea beneficiarului
33600000-6	Total lot 7					6385,00	7662,00	
	TOTAL					6385,00	7662,00	
						49240,00	58188,00	

Semnat:

Numele, prenumele: Ceaicovschi Tudor În calitate de: Director general

Ofertantul: „GBG-MLD” SRL

Adresa: mun. Chișinău, str. Tighina, 65, of. 607
 Pagina 1 din 1

REPUBLICA



MOLDOVA

CERTIFICAT DE ÎNREGISTRARE

PRIN PREZENTUL SE CERTIFICĂ, CĂ ÎNTRERINDEREAA
MIXTĂ "GBG-MLD" S.R.L. ESTE ÎNREGISTRATĂ LA CAMERA
ÎNREGISTRĂRII DE STAT

Numărul de indentificare de stat - codul fiscal

1003600117582

Data înregistrării

06.01.1995

Data eliberării

21.12.2004

Iovu Galina, registrator de stat

Funcția, numele, prenumele persoanei
care a eliberat certificatul

G. Iovu
semnătura

MD 0006733





AGENȚIA SERVICII PUBLICE

Departamentul înregistrare și licențiere a unităților de drept

EXTRAS din Registrul de stat al persoanelor juridice

Nr. 399048 data 03.12.2018

Denumirea completă: **Societatea cu Răspundere Limitată "GBG-MLD"**

Denumirea prescurtată: **"GBG-MLD" S.R.L.**

Forma juridică de organizare: **Societate cu răspundere limitată,**

Numărul de identificare de stat și codul fiscal (IDNO): **1003600117582**

Data înregistrării de stat: **06.01.1995**

Modul de constituire: **nou creată.**

Sediul: **MD-2001, str. Tighina, 65, mun. Chișinău, Republica Moldova.**

Obiectul principal de activitate:

1. **Comerțul cu ridicata al produselor farmaceutice**
2. **Cercetare și dezvoltare în științe fizice și naturale**
3. **Comerțul cu amănuntul al produselor farmaceutice și de parfumerie**
4. **Producția echipamentului de control pentru procesele industriale**
5. **Practica medicală**
6. **Fabricarea utilajului medical și chirurgical și a dispozitivelor ortopedice**
7. **Producția de aparatură și instrumente de măsură, verificare și control**
8. **Transporturi rutiere de mărfuri**

Capitalul social: **5400 lei,**

Administrator: **CEAICOVSCHI TUDOR, IDNP 0971601546960**

Asociații:

1. **COLEVA VERA, IDNP 2000048101473, cota 108 lei, ce constituie 2%**

2. **CEAICOVSCHI TUDOR, IDNP 0971601546960, cota 5292 lei, ce constituie 98%**

Beneficiar efectiv:

2.1. **CEAICOVSCHI TUDOR, IDNP 0971601546960, cota - 98%**

Prezentul extras este eliberat în temeiul art.34 al Legii nr.220-XVI din 19 octombrie 2007 privind înregistrarea de stat a persoanelor juridice și a întreprinzătorilor individuali și confirmă datele din Registrul de stat la data de: **03.12.2018.**

Registrator



Lozovanu Constantin
Lozovanu Constantin



EB 0249571



Certificate

The Certification Body of
TÜV Rheinland LGA Products GmbH

Hereby certifies that the organization

EUROIIMMUN
Medizinische Labordiagnostika AG
Seekamp 31
23560 Lübeck
Deutschland

has established and applies a quality management system for medical devices
for the following scope:

see attachment

Proof has been furnished that the requirements specified in

EN ISO 13485:2016

are fulfilled. The quality management system is subject to yearly surveillance

Effective Date: 2018-06-08

Certificate Registration No.: SX 60129534 0001

An audit was performed. Report No.: 21264033 005

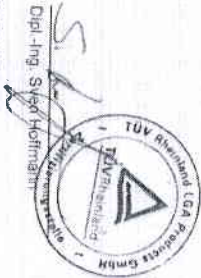
This Certificate is valid until: 2020-05-18

Certification Body



Date: 2018-06-08

TÜV Rheinland LGA Products GmbH - Tillystraße 2 - 90431 Nürnberg
Tel: +49 201 906-1371 Fax: +49 201 906-2000 Email: cert@tuev-rheinland.com Web: www.tuev-rheinland.com



Doc. 1/3, Rev. 0

TÜV Rheinland
LGA Products GmbH
Tillystraße 2, 90431 Nürnberg

Attachment to Certificate
Registration No.: SX 60129534 0001
Report No.: 21264033 005

Organization:
EUROIIMMUN
Medizinische Labordiagnostika AG
Seekamp 31
23560 Lübeck
Deutschland

Scope:

Design and development, production, installation, service and distribution of immunohistochemical test systems, immuno-fluorescence test systems, molecular diagnostic/genetic test systems, test systems for the determination of infectious agents and instruments/software for in vitro diagnostic

Sites included:

EUROIIMMUN Medizinische Labordiagnostika AG
Marktstraße 2-22, 23542 Dannew, Germany

Activities: Design and development, production, distribution

Certification Body



Date: 2018-06-08

Dipl.-Ing. Sven Hoffmann





Doc. 2/3, Rev. 0

TÜV Rheinland
LGA Products GmbH
Tillystraße 2, 90431 Nürnberg

Attachment to
Certificate
Registration No.: SX 60129534 0001
Report No.: 21264033 005

Organization:
EUROIMMUN
Medizinische Labordiagnostika AG
Seekamp 31
23560 Lübeck
Deutschland

Scope:

Sites included:

EUROIMMUN Medizinische Labordiagnostika AG
Am Sonnenberg 3/ 23627 Groß Gerau, Germany
Activities: Design and development, production
EUROIMMUN Medizinische Labordiagnostika AG
Am Born 24, 23627 Groß Gerau, Germany
Activities: Design and development, distribution,
installation, service
EUROIMMUN Medizinische Labordiagnostika AG
Im Kiepel 1, 92747 Heunburg, Germany
Activities: Production

Certification Body



Deutsche
Akkreditierungsstelle
D-20147 Hamburg

Date: 2018-06-08

S. Hoffmann
Dipl.-Ing. Sven Hoffmann



Doc. 2/3, Rev. 0

TÜV Rheinland
LGA Products GmbH
Tillystraße 2, 90431 Nürnberg

Attachment to
Certificate
Registration No.: SX 60129534 0001
Report No.: 21264033 005

Organization:
EUROIMMUN
Medizinische Labordiagnostika AG
Seekamp 31
23560 Lübeck
Deutschland

Scope:

Sites included:

EUROIMMUN Medizinische Labordiagnostika AG
Am Pleßhitzer 1, 92748 Bernsdorf, Germany
Activities: Production
EUROIMMUN Medizinische Labordiagnostika AG
Schloßstraße 11, 91257 Regnitz, Germany
Activities: Production, installation, service
EUROIMMUN Medizinische Labordiagnostika AG
Am der Traße 2, 23923 Selmsdorf, Germany
Activities: Design and development, production, service

Certification Body



Deutsche
Akkreditierungsstelle
D-20147 Hamburg

Date: 2018-06-08

S. Hoffmann
Dipl.-Ing. Sven Hoffmann





Anti-Measles Virus ELISA (IgG) Test instruction

ORDER NO.	ANTIBODIES AGAINST	IG CLASS	SUBSTRATE	FORMAT
EI 2610-9601 G	Measles virus	IgG	Ag-coated microplate-wells	96 x 01 (96)

Indication: measles

Principles of the test: The ELISA test kit provides a quantitative or semiquantitative in vitro assay for human antibodies of the IgG class against measles virus in serum or plasma. The test kit contains microtiter strips each with 8 break-off reagent wells coated with measles virus antigens. In the first reaction step, diluted patient samples are incubated in the wells. In the case of positive samples, specific IgG antibodies (also IgA and IgM) will bind to the antigens. To detect the bound antibodies, a second incubation is carried out using an enzyme-labelled anti-human IgG (enzyme conjugate) catalysing a colour reaction.

Contents of the test kit:

Component	Colour	Format	Symbol
1. Microplate wells coated with antigens: 12 microplate strips each containing 8 individual break-off wells in a frame, ready for use	---	12 x 8	[STRIPS]
2. Calibrator 1 5000 IU/l (IgG, human), ready for use		1 x 2.0 ml	[CAL 1]
3. Calibrator 2 1000 IU/l (IgG, human), ready for use	red coloured in decreasing intensity	1 x 2.0 ml	[CAL 2]
4. Calibrator 3 250 IU/l (IgG, human), ready for use		1 x 2.0 ml	[CAL 3]
5. Calibrator 4 50 IU/l (IgG, human), ready for use		1 x 2.0 ml	[CAL 4]
6. Positive control (IgG, human), ready for use	blue	1 x 2.0 ml	[POS CONTROL]
7. Negative control (IgG, human), ready for use	green	1 x 2.0 ml	[NEG CONTROL]
8. Enzyme conjugate peroxidase-labelled anti-human IgG (rabbit), ready for use	green	1 x 12 ml	[CONJUGATE]
9. Sample buffer ready for use	light blue	1 x 100 ml	[SAMPLE BUFFER]
10. Wash buffer 10x concentrate	colourless	1 x 100 ml	[WASH BUFFER 10X]
11. Chromogen/substrate solution TMB/H ₂ O ₂ , ready for use	colourless	1 x 12 ml	[SUBSTRATE]
12. Stop solution 0.5 M sulphuric acid, ready for use	colourless	1 x 12 ml	[STOP SOLUTION]
13. Test instruction		1 booklet	
14. Quality control certificate		1 protocol	
LOT Lot	CE	Storage temperature	
IVD In vitro determination		Unopened usable until:	

Storage and stability: The test kit has to be stored at a temperature between +2°C to +8°C. Do not freeze. Unopened, all test kit components are stable until the indicated expiry date.

Waste disposal: Patient samples, calibrators, controls and incubated microplate strips should be handled as infectious waste. All reagents must be disposed of in accordance with local disposal regulations.

Preparation and stability of the reagents

Note: All reagents must be brought to room temperature (+18°C to +25°C) approx. 30 minutes before use. After first use, the reagents are stable until the indicated expiry date if stored at +2°C to +8°C and protected from contamination, unless stated otherwise below.

Coated wells: Ready for use. Tear open the resealable protective wrapping of the microplate at the recesses above the grip seam. Do not open until the microplate has reached room temperature to prevent the individual strips from moistening. Immediately replace the remaining wells of a partly used microplate in the protective wrapping and tightly seal with the integrated grip seam (Do not remove the desiccant bag).

Once the protective wrapping has been opened for the first time, the wells coated with antigens can be stored in a dry place and at a temperature between +2°C and +8°C for 4 months.

Calibrators and controls: Ready for use. The reagents must be mixed thoroughly before use.

Enzyme conjugate: Ready for use. The enzyme conjugate must be mixed thoroughly before use.

Sample buffer: Ready for use.

Wash buffer: The wash buffer is a 10x concentrate. If crystallization occurs in the concentrated buffer, warm it to 37°C and mix well before diluting. The quantity required should be removed from the bottle using a clean pipette and diluted with deionized or distilled water (1 part reagent plus 9 parts distilled water).

For example: For 1 microplate strip, 5 ml concentrate plus 45 ml water.
The working strength wash buffer is stable for 4 weeks when stored at +2°C to +8°C and handled properly.

Chromogen/substrate solution: Ready for use. Close the bottle immediately after use, as the contents are sensitive to light. The Chromogen/substrate solution must be clear on use. Do not use the solution if it is blue coloured.

Stop solution: Ready for use.

Warning: The controls and calibrators used have been tested negative for HBSAg, anti-HCV, anti-HIV-1 and anti-HIV-2 using enzyme immunoassays and indirect immunofluorescence methods. Nonetheless, all materials should be treated as being a potential infection hazard and should be handled with care. Some of the reagents contain the toxic agent sodium azide. Avoid skin contact.

Preparation and stability of the patient samples

Samples: Human serum or EDTA, heparin or citrate plasma.

Stability: Patient samples to be investigated can generally be stored at +2°C to +8°C for up to 14 days. Diluted samples should be incubated within one working day.

Sample dilution: Patient samples are diluted 1:101 in sample buffer. For example: dilute 10 µl serum in 1.0 ml sample buffer and mix well by vortexing (sample pipettes are not suitable for mixing).

NOTE: Calibrators and controls are prediluted and ready for use, do not dilute them.



Incubation

For semiquantitative analysis: incubate calibrator 3 along with the positive and negative controls and patient samples. For quantitative analysis: incubate calibrators 1 to 4 along with the positive and negative controls and patient samples.

(Partly) manual test performance

Sample incubation: (1st step)

Transfer 100 µl of the calibrators, positive and negative controls or diluted patient samples into the individual microplate wells according to the pipetting protocol. Incubate for 30 minutes at room temperature (+18°C to +25°C).

Washing:

Manual: Empty the wells and subsequently wash 3 times using 300 µl of working strength wash buffer for each wash.
Automatic: Wash reagent wells 3 times with 450 µl of working strength wash buffer (program setting: e.g. TECAN Columbus Washer "Overflow Modus").

Leave the wash buffer in each well for 30 to 60 seconds per washing cycle, then empty the wells. After washing (manual and automated tests), thoroughly dispose of all liquid from the microplate by tapping it on absorbent paper with the openings facing downwards to remove all residual wash buffer.

Note: Residual liquid (> 10 µl) in the reagent wells after washing can interfere with the substrate and lead to false low extinction values. Insufficient washing (e.g., less than 3 wash cycles, too small wash buffer volumes, or too short reaction times) can lead to false high extinction values.
Free positions on the microplate strip should be filled with blank wells of the same plate format as that of the parameter to be investigated.

Conjugate incubation: (2nd step)

Pipette 100 µl of enzyme conjugate (peroxidase-labelled anti-human IgG) into each of the microplate wells. Incubate for 30 minutes at room temperature (+18°C to +25°C).

Washing:

Empty the wells. Wash as described above.

Substrate incubation: (3rd step)

Pipette 100 µl of chromogen/substrate solution into each of the microplate wells. Incubate for 15 minutes at room temperature (+18°C to +25°C) (protect from direct sunlight).

Stopping the reaction:

Pipette 100 µl of stop solution into each of the microplate wells in the same order and at the same speed as the chromogen/substrate solution was introduced.

Measurement:

Photometric measurement of the colour intensity should be made at a wavelength of 450 nm and a reference wavelength between 620 nm and 650 nm within 30 minutes of adding the stop solution. Prior to measuring, slightly shake the microplate to ensure a homogeneous distribution of the solution.

Test performance using fully automated analysis devices

Sample dilution and test performance are carried out fully automatically using the analysis device. The incubation conditions programmed in the respective software authorised by EUROIMMUN may deviate slightly from the specifications given in the ELISA test instruction. However, these conditions were validated in respect of the combination of the EUROIMMUN Analyzer I or the DSX from Dynex and this EUROIMMUN ELISA. Validation documents are available on inquiry.
Automated test performance using other fully automated, open system analysis devices is possible, however, the combination should be validated by the user.



Pipetting protocol

	1	2	3	4	5	6	7	8	9	10	11	12
A	C3	P 6	P 14	P 22			C1	P 3	P 11	P 19		
B	pos.	P 7	P 15	P 23			C2	P 4	P 12	P 20		
C	neg.	P 8	P 16	P 24			C3	P 5	P 13	P 21		
D	P 1	P 9	P 17				C4	P 6	P 14	P 22		
E	P 2	P 10	P 18				pos.	P 7	P 15	P 23		
F	P 3	P 11	P 19				neg.	P 8	P 16	P 24		
G	P 4	P 12	P 20				P 1	P 9	P 17			
H	P 5	P 13	P 21				P 2	P 10	P 18			

The pipetting protocol for microtiter strips 1-4 is an example for the semiquantitative analysis of 24 patient samples (P 1 to P 24).

The pipetting protocol for microtiter strips 7-10 is an example for the quantitative analysis of 24 patient samples (P 1 to P 24).

The calibrators (C 1 to C 4), the positive (pos.) and negative (neg.) controls, and the patient samples have each been incubated in one well. The reliability of the ELISA test can be improved by duplicate determinations for each sample.

The reagent wells are break-off format. Therefore, the number of tests performed can be matched to the number of samples, minimizing reagent wastage.

Both positive and negative controls serve as internal controls for the reliability of the test procedure. They should be assayed with each test run.

Calculation of results

Semiquantitative: Results can be evaluated semiquantitatively by calculating a ratio of the extinction value of the control or patient sample over the extinction value of calibrator 3. Calculate the ratio according to the following formula:

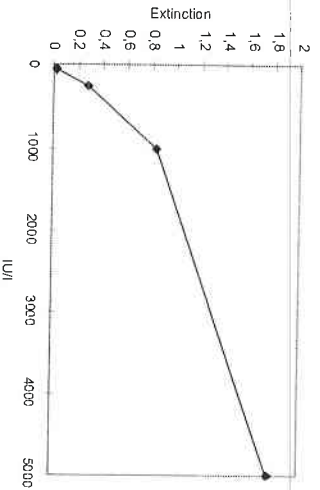
$$\frac{\text{Extinction of the control or patient sample}}{\text{Extinction of calibrator 3}} = \text{Ratio}$$

EUROIMMUN recommends interpreting results as follows:

Ratio < 0.8:	negative
Ratio ≥ 0.8 to < 1.1:	borderline
Ratio ≥ 1.1:	positive

In cases of borderline test results, an additional patient sample should be taken 7 days later and re-tested in parallel with the first patient sample. The results of both samples allow proper evaluation of titer changes.

Quantitative: The standard curve from which the concentration of antibodies in the patient samples can be taken is obtained by point-to-point plotting of the extinction values measured for the 4 calibrators against the corresponding units (linear/linear). Use "point-to-point" plotting for calculation of the standard curve by computer. The following plot is an example of a typical calibration curve. Please do not use this curve for the determination of antibody concentrations in patient samples.



If the extinction of a serum sample lies above the value of calibrator 1 (5000 IU/l), the result should be given as ">5000 IU/l". It is recommended that the sample be re-tested at a dilution of 1:400. The result in IU/l read from the calibration curve for this sample must then be multiplied by a factor of 4.

The upper limit of the reference range of non-infected persons (cut-off value) recommended by EUROIMMUN is 250 International Units (IU/l). EUROIMMUN recommends interpreting results as follows:

- <200 IU/l: negative
- ≥200 to <275 IU/l: borderline
- ≥275 IU/l: positive

Evaluation information: For duplicate determinations the mean of the two values should be taken. If the two values deviate substantially from one another the sample should be retested.

For the interpretation of borderline results an investigation using further tests (e.g. avidity determination of antibody class IgG) can be helpful. Diagnosis can be secured by the determination of the titer change in two serum samples taken at an interval of at least 7 days and analysed in parallel.

For diagnosis, the clinical symptoms of the patient should always be taken into account along with the serological results.

Test characteristics

Calibration: The controls of the Anti-Measles Virus ELISA (IgG) were calibrated using the 3rd international standard serum NIBSC 97/648 (anti-measles and anti-polio virus serum, National Institute for Biological Standards and Control, Hertfordshire, England, approved as international reference preparation by the WHO Expert Committee on Biological Standardization). The NIBSC 97/648 serum contains 3 International Units (IU) per ampoule by definition and was resuspended in a concentration of 3 IU/ml.

For every group of tests performed, the extinction values of the calibrators and the international units determined for the positive and negative controls must lie within the limits stated for the relevant test kit lot. A quality control certificate containing these reference values is included. If the values specified for the controls are not achieved, the test results may be inaccurate and the test should be repeated.

The activity of the enzyme used is temperature-dependent and the extinction values may vary if a thermostat is not used. The higher the room temperature during substrate incubation, the greater will be the extinction values. Corresponding variations apply also to the incubation times. However, the calibrators are subject to the same influences, with the result that such variations will be largely compensated in the calculation of the result.

Antigen: The antigen source is provided by inactivated cell lysates of Vero cells infected with the "Edmonston" strain of measles viruses.

Linearity: The linearity of the anti-measles viruses ELISA (IgG) was determined by assaying 4 serial dilutions of different patient samples. The coefficient of determination R^2 for all sera was > 0.95. The Anti-Measles Virus ELISA (IgG) is linear at least in the tested concentration range (52 IU/l - 4865 IU/l).

Detection limit: The lower detection limit is defined as the mean value of an analyte-free sample plus three times the standard deviation and is the smallest detectable antibody titer. The lower detection limit of the Anti-Measles Virus ELISA (IgG) is 8 IU/l.

Cross reactivity: The quality of the antigen used ensures a high specificity of the ELISA. Sera from patients with infections caused by various agents were investigated with the EUROIMMUN Anti-measles virus ELISA (IgG)

Antibodies against	n	Anti-Measles virus ELISA (IgG)
Adenovirus	8	0%
CMV	6	0%
EBV/CA	11	0%
HSV-1	3	0%
Influenza virus type A	5	0%
Influenza virus type B	11	0%
Mumps virus	4	0%
Mycoplasma pneumoniae	4	0%
Parainfluenza virus types 1-4	11	0%
RSV	9	0%
Rubella virus	6	0%
Toxoplasma	3	0%
VZV	5	0%

Interference: Haemolytic, lipaemic and icteric samples showed no influence on the result up to a concentration of 10 mg/ml for hemoglobin, 20 mg/ml for triglycerides and 0.4 mg/ml for bilirubin in this ELISA.

Reproducibility: The reproducibility of the test was investigated by determining the intra- and inter-assay coefficients of variation using 3 sera. The intra-assay CVs are based on 20 determinations and the inter-assay CVs on 4 determinations performed in 6 different test runs.

Serum	Intra-assay variation, n = 20	
	Mean value (IU/l)	CV (%)
1	830	8.0
2	3410	6.6
3	3725	5.6

Serum	Inter-assay variation, n = 4 x 6	
	Mean value (IU/l)	CV (%)
1	796	11.6
2	3635	5.0
3	3946	6.8



Specificity and sensitivity: 112 clinically characterized patient samples (interlaboratory test samples from INSTAND, Germany, Labquality, Finland and NEQAS, UK) were examined with the EUROIMMUN Anti-Measles Virus ELISA (IgG). The test showed a specificity and a sensitivity of 100% each.

n = 112	INSTAND / Labquality / NEQAS (IgG)			
	positive	borderline	negative	
EUROIMMUN	89	1	0	
Anti-Measles Virus	borderline	0	0	
ELISA (IgG)	negative	0	0	22

Reference range: The levels of anti-measles virus antibodies (IgG) were analyzed with this EUROIMMUN ELISA in a panel of 500 healthy blood donors. With a cut-off of 250 IU/l, 94% of the blood donors were anti-measles virus positive (IgG), which reflects the known percentage of infections in adults.

Clinical significance

The measles virus (MV) is the most instantly recognizable member of Morbilliviruses, a group of viruses belonging to the Paramyxoviridae family [1]. No animal reservoir is known. The measles virus causes an acute febrile illness which occurs mainly in childhood and is very infectious [2, 3]. In 1999, measles still caused worldwide 873,000 deaths per year [1, 4, 5]. Today they are less frequent because of vaccination, especially in the western hemisphere [6, 7]. However, measles epidemics are still observed in some countries [2, 3, 4, 5, 6, 8, 9]. Individuals acutely infected with the virus exhibit a wide range of clinical symptoms ranging from a characteristic mild self-limiting infection to death [1, 2, 8, 10].

MV infections are characterised by an incubation period of about 10 days. flu-like symptoms with fever, malaise, catarrh of the upper respiratory tract, cough, congestion and conjunctivitis. Soon afterwards the measles rash, a typical exanthema, appears first near the ears, then on the forehead, in the face and over the rest of the body [1, 5, 8, 11].

Complications arising from MV infections include secondary bacterial pneumonia, otitis media (approx. 1%), encephalitis (approx. 1%), myocarditis, miscarriage and a condition called subacute sclerosing panencephalitis (SSPE) [5, 12, 13]. Persistent MV infection of the otic capsule is an aetiological factor in otosclerosis [9, 14]. Anti-measles IgG for the serological diagnosis of otosclerotic hearing loss has a high specificity and sensitivity [15]. SSPE is a progressive, generally fatal brain disorder caused by chronic measles virus infection. It occurs about 7 to 10 years after the infection and generally kills within 3 years from the onset of the symptoms. The patients suffer from behavioural changes, cognitive deterioration, vision problems and eventually advanced neurological symptoms, such as severe spasms, and finally severe physical and mental impairment that leads to death [13]. Males are more commonly affected than females. The risk of SSPE from measles was underestimated according to older data [12]. Actual papers put it at closer to 6.5 to 11 cases of SSPE per 100,000 measles infections; that means 7 to 13 times higher than the earlier estimates [1, 9, 12].

Women with acute measles infection during pregnancy and a negative result for measles-specific antibodies were observed e.g. in Japan, India, Thailand, Kenya and Brazil: 3 of 4 pregnancies ended in preterm delivery, spontaneous abortion or stillbirth, 2 of 4 neonates were found to have congenital measles with a positive result for IgM antibodies [5].

Antibodies against MV can be found in the serum of almost all patients during and after a measles infection. IgM antibodies develop soon after the onset of symptoms and can be measured using ELISA or indirect immunofluorescence tests (IFIT) [16, 20, 21, 22]. 50% of patients have IgM antibodies within three days, more than 90% within 10 days after occurrence of the rash [15, 17]. The Anti-Measles Virus IgM ELISA is more rapid and sensitive for the serological diagnosis of measles infections than other tests [15, 18, 19].



MV infections often cause an increase in heterologic antibodies. The statistically evaluated detection rate for antibodies is significantly higher for ELISA and IFIT in comparison with e.g. neutralisation tests [16, 20]. IgG and IgM antibodies against MV are reliable markers to confirm suspected measles infections.

Measles myelitis or encephalitis can be verified by detecting antibodies against measles in the cerebrospinal fluid (CSF) [23, 24, 25, 26]. These specific antibodies are synthesised in the brain [24]. The CSF-serum quotient (CSQ) allows to differentiate between a blood-derived and a pathological, brain-derived specific antibody fraction in CSF, taking into account individual changes in the blood/CSF barrier function [24, 25, 26, 27]. Therefore it is necessary to confirm the presence of antibodies against MV using ELISA both in CSF and in the serum. During measles myelitis or encephalitis an intrathecal synthesis of antibodies against MV in CSF takes place. Due to the fact that specific antibodies can pass the blood-cerebrospinal fluid barrier by diffusion from serum to CSF it is necessary to determine the relative CSF/serum quotient (CSQrel, synonym: antibody specificity index) [24, 25, 26]. The quotient is calculated from the amount of specific anti-measles virus IgG antibodies in total CSF IgG in proportion to the amount of specific IgG antibodies in total serum IgG. During conversion the CSF/serum quotient of the pathogen-specific IgG-antibody concentrations CSQpath-spec. (IgG) is put into relation to the CSF/serum quotient of the total IgG concentrations CSQtotal (IgG) [27]. A relative CSQ result above 1.5 indicates the production of specific antibodies in the central nervous system (CNS) and the involvement of the CNS in the disease [25, 26].

With respect to the severe complications known from measles infections, the Robert Koch Institute in Germany recommends vaccinating small children, with a first shot between the age of 11 to 14 months and a second between 15 and 23 months [2, 4, 10, 11]. Neutralisation activity and persistence of antibodies are induced in response to the immunisation [6, 15].

Life-long immunity is generally developed. However, antibody levels are 8 to 10 times lower in post-vaccination sera than in convalescent sera [6, 19, 28]. A passive immunisation with specific immunoglobulin concentrates is usually given to immunosuppressed seronegative individuals, such as tumour patients and recipients of transplants, as well as to seronegative pregnant women after exposure to the virus.

The European Regional Office of the WHO aims at eliminating measles from the region in the following years by area-wide vaccination campaigns [4, 9, 10]. This is expected to limit the number of apparent infections and especially of severe courses of the disease. For the diagnosis of the remaining cases of measles infection and of infections acquired outside Europe as well as for the clarification of atypical courses of the disease in partly immunised patients the antibody determination in serum and CSF will be of growing importance [3, 5, 8, 9].



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Anti-Measles Virus ELISA (IgM) Test instruction

ORDER NO.	ANTIBODIES AGAINST	IG CLASS	SUBSTRATE	FORMAT
El 2610-9601 M	Measles virus	IgM	Ag-coated microplate wells	96 x 01 (96)

Principles of the test: The ELISA test kit provides a semiquantitative in vitro assay for human antibodies of the IgM class against measles virus in serum or plasma. The test kit contains microtiter strips each with 8 break-off reagent wells coated with measles virus antigens. In the first reaction step, diluted patient samples are incubated in the wells. In the case of positive samples, specific IgM antibodies (also IgA and IgG) will bind to the antigens. To detect the bound antibodies, a second incubation is carried out using an enzyme-labelled anti-human IgM (enzyme conjugate) catalysing a colour reaction.

Contents of the test kit:

Component	Colour	Format	Symbol
1. Microplate wells coated with antigens: 12 microplate strips each containing 8 individual break-off wells in a frame, ready for use	---	12 x 8	[STRIPS]
2. Calibrator (IgM, human), ready for use	dark red	1 x 20 ml	[CAL]
3. Positive control (IgM, human), ready for use	blue	1 x 20 ml	[POS CONTROL]
4. Negative control (IgM, human), ready for use	green	1 x 20 ml	[NEG CONTROL]
5. Enzyme conjugate peroxidase-labelled anti-human IgM (goat), ready for use	red	1 x 12 ml	[CONJUGATE]
6. Sample buffer containing IgG/RF absorbent (anti-human IgG antibody preparation obtained from goat), ready for use	green	1 x 100 ml	[SAMPLE BUFFER]
7. Wash buffer 10x concentrate	colourless	1 x 100 ml	[WASH BUFFER 10X]
8. Chromogen/substrate solution TMB/H ₂ O ₂ , ready for use	colourless	1 x 12 ml	[SUBSTRATE]
9. Stop solution 0.5 M sulphuric acid, ready for use	colourless	1 x 12 ml	[STOP SOLUTION]
10. Test instruction	---	1 booklet	
11. Quality control certificate	---	1 protocol	
LOT In vitro determination			
IVD			

Storage and stability: The test kit has to be stored at a temperature between +2°C to +8°C. Do not freeze. Unopened, all test kit components are stable until the indicated expiry date.

Waste disposal: Patient samples, calibrators, controls and incubated microplate strips should be handled as infectious waste. All reagents must be disposed of in accordance with local disposal regulations.

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Preparation and stability of the reagents

Note: All reagents must be brought to room temperature (+18°C to +25°C), approx. 30 minutes before use. After first use, the reagents are stable until the indicated expiry date if stored at +2°C to +8°C and protected from contamination, unless stated otherwise below.

Coated wells: Ready for use. Tear open the resealable protective wrapping of the microplate at the recesses above the grip seam. Do not open until the microplate has reached room temperature to prevent the individual strips from moistening. Immediately replace the remaining wells of a partly used microplate in the protective wrapping and tightly seal with the integrated grip seam (Do not remove the desiccant bag).
Once the protective wrapping has been opened for the first time, the wells coated with antigens can be stored in a dry place and at a temperature between +2°C and +8°C for 4 months.

Calibrator and controls: Ready for use. The reagents must be mixed thoroughly before use.

Enzyme conjugate: Ready for use. The enzyme conjugate must be mixed thoroughly before use.

Sample buffer: Ready for use. The green coloured sample buffer contains IgG/RF absorbent. Serum or plasma samples diluted with this sample buffer are only to be used for the determination of IgM antibodies.

Wash buffer: The wash buffer is a 10x concentrate. If crystallization occurs in the concentrated buffer, warm it to 37°C and mix well before diluting. The quantity required should be removed from the bottle using a clean pipette and diluted with deionized or distilled water (1 part reagent plus 9 parts distilled water).

For example: For 1 microplate strip, 5 ml concentrate plus 45 ml water.
The working strength wash buffer is stable for 4 weeks when stored at +2°C to +8°C and handled properly.

Chromogen/substrate solution: Ready for use. Close the bottle immediately after use, as the contents are sensitive to light. The Chromogen/substrate solution must be clear on use. Do not use the solution if it is blue coloured.

Stop solution: Ready for use.

Warning: The controls and calibrators used have been tested negative for HBsAg, anti-HCV, anti-HIV-1 and anti-HIV-2 using enzyme immunoassays and indirect immunofluorescence methods. Nonetheless, all materials should be treated as being a potential infection hazard and should be handled with care. Some of the reagents contain the toxic agent sodium azide. Avoid skin contact.



Preparation and stability of the patient samples

Samples: Human serum or EDTA, heparin or citrate plasma.

Stability: Patient samples to be investigated can generally be stored at +2°C to +8°C for up to 14 days. Diluted samples should be incubated within one working day.

Introduction: Before the determination of specific antibodies of class IgM, antibodies of class IgG should be removed from the patient sample. This procedure must be carried out in order to prevent any rheumatoid factors from reacting with specifically bound IgG, which would lead to false positive IgM test results, and to prevent specific IgG displacing IgM from the antigen, which would lead to false IgM negative test results.

Functional principle: The sample buffer (green coloured) contains an anti-human antibody preparation from goat IgG from a serum sample is bound with high specificity by these antibodies and precipitated. If the sample also contains rheumatoid factors, these will be absorbed by the IgG/anti-human IgG complex.

Separation properties:

- All IgG subclasses are bound and precipitated by the anti-human IgG antibodies.
- Human serum IgG in concentrations of up to 15 mg per ml are removed (average serum IgG concentration in adults: 12 mg per ml).
- Rheumatoid factors are also removed.
- The recovery rate of the IgM fraction is almost 100%.

Performance: The patient samples for analysis are diluted 1:101 with sample buffer. For example, add 10 µl sample to 1.0 ml sample buffer and mix well. Incubate the mixture for at least 10 minutes at room temperature. Subsequently, it can be pipetted into the microplate wells according to the pipetting protocol.

Notes:

- Antibodies of the class IgG should not be analyzed with this mixture.
 - It is possible to check the efficacy of the IgG/RF absorbent for an individual patient sample by performing an IgG test in parallel to the IgM test using the mixture. If the IgG test is negative, the IgM result can be considered as reliable.
- The calibrator and controls containing IgM antibodies are pre-diluted and ready for use, do not dilute them



Incubation

(Partly) manual test performance

Sample incubation:
(1st step)

Transfer 100 µl of the calibrator, positive and negative controls or diluted patient samples into the individual microplate wells according to the pipetting protocol. Incubate for 30 minutes at room temperature (+18°C to +25°C).

Washing:

Manual: Empty the wells and subsequently wash 3 times using 300 µl of working strength wash buffer for each wash.

Automatic: Wash reagent wells 3 times with 450 µl working strength wash buffer (program setting: e.g. TECAN Columbus Washer "Overflow Modus").

Leave the wash buffer in each well for 30 to 60 seconds per washing cycle, then empty the wells. After washing (manual and automated tests), thoroughly dispose of all liquid from the microplate by tapping it on absorbent paper with the openings facing downwards to remove all residual wash buffer.

Note: Residual liquid (> 10 µl) in the reagent wells after washing can interfere with the substrate and lead to false low extinction values. Insufficient washing (e.g., less than 3 wash cycles, too small wash buffer volumes, or too short reaction times) can lead to false high extinction values.

Free positions on the microplate strip should be filled with blank wells of the same plate format as that of the parameter to be investigated.

Conjugate incubation:
(2nd step)

Pipette 100 µl of enzyme conjugate (peroxidase-labelled anti-human IgM) into each of the microplate wells. Incubate for 30 minutes at room temperature (+18°C to +25°C).

Washing:

Empty the wells. Wash as described above.

Substrate incubation:
(3rd step)

Pipette 100 µl of chromogen/substrate solution into each of the microplate wells. Incubate for 15 minutes at room temperature (+18°C to +25°C) (protect from direct sunlight).

Stopping the reaction:

Pipette 100 µl of stop solution into each of the microplate wells in the same order and at the same speed as the chromogen/substrate solution was introduced.

Measurement:

Photometric measurement of the colour intensity should be made at a wavelength of 450 nm and a reference wavelength between 620 nm and 650 nm within 30 minutes of adding the stop solution. Prior to measuring, slightly shake the microplate to ensure a homogeneous distribution of the solution.

Test performance using fully automated analysis devices

Sample dilution and test performance are carried out fully automatically using the analysis device. The incubation conditions programmed in the respective software authorised by EUROIMMUN may deviate slightly from the specifications given in the ELISA test instruction. However, these conditions were validated in respect of the combination of the EUROIMMUN Analyzer I, Analyzer I-2P or the DSX from Dynex and this EUROIMMUN ELISA. Validation documents are available on inquiry. Automated test performance using other fully automated, open system analysis devices is possible, however, the combination should be validated by the user.



Pipetting protocol

1	2	3	4	5	6	7	8	9	10	11	12
A	C	P 6	P 14	P 22							
B	P 5a	P 7	P 15	P 23							
C	neg	P 8	P 16	P 24							
D	P 1	P 9	P 17								
E	P 2	P 10	P 18								
F	P 3	P 11	P 19								
G	P 4	P 12	P 20								
H	P 5	P 13	P 21								

The above pipetting protocol is an example of the semiquantitative analysis of antibodies in 24 patient samples (P 1 to P 24).

Calibrator (C), positive (pos.) and negative (neg.) control as well as the patient samples have been incubated in one well each. The reliability of the ELISA test can be improved by duplicate determinations of each sample.

The wells can be broken off individually from the strips. This makes it possible to adjust the number of test substrates used to the number of samples to be examined and minimizes reagent wastage. Both positive and negative controls serve as internal controls for the reliability of the test procedure. They should be assayed with each test run.

Calculation of results

The extinction value of the calibrator defines the upper limit of the reference range of non-infected persons (cut-off) recommended by EUROIMMUN. Values above the indicated cut-off are to be considered as positive, those below as negative.

Semiquantitative: Results can be evaluated semiquantitatively by calculating a ratio of the extinction value of the control or patient sample over the extinction value of calibrator. Use the following formula to calculate the ratio:

$$\frac{\text{Extinction of the control or patient sample}}{\text{Extinction of calibrator}} = \text{Ratio}$$

EUROIMMUN recommends interpreting results as follows:

- Ratio < 0.8: negative
- Ratio ≥ 0.8 to < 1.1: borderline
- Ratio ≥ 1.1: positive

Evaluation information: For duplicate determinations the mean of the two values should be taken. If the two values deviate substantially from one another the sample should be retested.

For the interpretation of borderline results an investigation using further tests (e.g. avidity determination of antibody class (IgG) can be helpful. Diagnosis can be secured by the determination of the titer change in two serum samples taken at an interval of at least 7 days and analysed in parallel.

For diagnosis, the clinical symptoms of the patient should always be taken into account along with the serological results.



Test characteristics

Calibration: As no international reference serum exists for antibodies of the IgM class against measles virus, results are provided in the form of ratios which are a relative measure for the concentration of antibodies.

For every group of tests performed, the extinction values of the calibrator and the ratios of the positive and negative controls must lie within the limits stated for the relevant test kit lot. A quality control certificate containing these reference values is included. If the values specified for the controls are not achieved, the test results may be inaccurate and the test should be repeated.

The activity of the enzyme used is temperature-dependent and the extinction values may vary if a thermostat is not used. The higher the room temperature during substrate incubation, the greater will be the extinction values. Corresponding variations apply also to the incubation times. However, the calibrators are subject to the same influences, with the result that such variations will be largely compensated in the calculation of the result.

Antigen: The antigen source is provided by inactivated cell lysates of Vero cells infected with the "Edmonston" strain of measles viruses.

Detection limit: The lower detection limit is defined as the mean value of an analyte-free sample plus three times the standard deviation and is the smallest detectable antibody titer. The lower detection limit of the Anti-Measles Virus ELISA (IgM) is ratio 0.02.

Cross reactivity: The quality of the antigen used ensures a high specificity of the ELISA. Sera from patients with infections caused by various agents were investigated with the EUROIMMUN Anti-Measles Virus ELISA (IgM).

Antibodies against	n	Anti-Measles Virus ELISA (IgM)
Borrelia burgdorferi	10	0%
CMV	7	0%
EBV CA	17	0%
Mumps virus	8	0%
Parvovirus B19	9	0%
Rubella virus	10	0%
Toxoplasma gondii	10	0%
VZV	5	0%

Interference: Haemolytic, lipaemic and icteric samples showed no influence on the result up to a concentration of 10 mg/ml for hemoglobin, 20 mg/ml for triglycerides and 0.4 mg/ml for bilirubin in this ELISA.

Reproducibility: The reproducibility of the test was investigated by determining the intra- and inter-assay coefficients of variation using 3 sera. The intra-assay CVs are based on 20 determinations and the inter-assay CVs on 4 determinations performed in 6 different test runs.

Serum	Mean value (Ratio)	CV (%)
1	2.6	7.9
2	4.6	2.5
3	7.0	2.3

Serum	Mean value (Ratio)	CV (%)
1	2.4	8.0
2	4.1	4.4
3	6.6	4.4



Specificity and sensitivity: 72 clinically characterized patient samples (interlaboratory test samples from INSTAND, Germany) were examined with the EUROIMMUN Anti-Measles Virus ELISA (IgM). The test showed a specificity of 98% and a sensitivity of 100%.

ELISA	INSTAND	
	positive	negative
positive	26	1
cut-off	0	1
negative	0	44

n = 72

Reference range: The levels of the anti-measles virus antibodies (IgM) were analyzed with this EUROIMMUN ELISA in a panel of 300 healthy blood donors. With a cut-off ratio of 1.0, 0.3% of the blood donors were anti-measles viruses positive (IgM).

Clinical significance

The measles virus (MV) is the most instantly recognizable member of Morbilliviruses, a group of viruses belonging to the Paramyxoviridae family [1]. No animal reservoir is known. The measles virus causes an acute febrile illness which occurs mainly in childhood and is very infectious [2, 3]. In 1999, measles still caused worldwide 873,000 deaths per year [1, 4, 5]. Today they are less frequent because of vaccination, especially in the western hemisphere [6, 7]. However, measles epidemics are still observed in some countries [2, 3, 4, 5, 6, 8, 9]. Individuals acutely infected with the virus exhibit a wide range of clinical symptoms ranging from a characteristic mild self-limiting infection to death [1, 2, 8, 10].

MV infections are characterized by an incubation period of about 10 days, flu-like symptoms with fever, malaise, catarrh of the upper respiratory tract, cough, congestion and conjunctivitis. Soon afterwards the measles rash, a typical exanthema, appears first near the ears, then on the forehead, in the face and over the rest of the body [1, 5, 8, 11].

Complications arising from MV infections include secondary bacterial pneumonia, otitis media (approx. 1%), encephalitis (approx. 1%), myocarditis, miscarriage and a condition called subacute sclerosing panencephalitis (SSPE) [5, 12, 13]. Persistent MV infection of the otic capsule is an etiologic factor in otosclerosis [9, 14]. Anti-measles IgG for the serological diagnosis of otosclerotic hearing loss has a high specificity and sensitivity [15]. SSPE is a progressive, generally fatal brain disorder caused by a chronic measles virus infection. It occurs about 7 to 10 years after the infection and generally kills within 3 years from the onset of the symptoms. The patients suffer from behavioural changes, cognitive deterioration, vision problems and eventually advanced neurological symptoms, such as severe spasms, and finally severe physical and mental impairment that leads to death [13]. Males are more commonly affected than females. The risk of SSPE from measles was underestimated according to older data [12]. Actual papers put it at closer to 6.5 to 11 cases of SSPE per 100,000 measles infections, that means 7 to 13 times higher than the earlier estimates [1, 9, 12].

Women with acute measles infection during pregnancy and a negative result for measles-specific antibodies were observed e.g. in Japan, India, Thailand, Kenya and Brazil: 3 of 4 pregnancies ended in preterm delivery, spontaneous abortion or stillbirth; 2 of 4 neonates were found to have congenital measles with a positive result for IgM antibodies [5].

Antibodies against MV can be found in the serum of almost all patients during and after a measles infection. IgM antibodies develop soon after the onset of symptoms and can be measured using ELISA or indirect immunofluorescence tests (IFIT) [16, 20, 21, 22]. 50% of patients have IgM antibodies within three days, more than 90% within 10 days after occurrence of the rash [15, 17]. The Anti-Measles Virus IgM ELISA is more rapid and sensitive for the serological diagnosis of measles infections than other tests [15, 18, 19]. MV infections often cause an increase in heterologic antibodies. The statistically evaluated detection rate for antibodies is significantly higher for ELISA and IFIT in comparison with e.g. neutralisation tests [16, 20]. IgG and IgM antibodies against MV are reliable markers to confirm suspected measles infections.



Measles myelitis or encephalitis can be verified by detecting antibodies against measles in the cerebrospinal fluid (CSF) [23, 24, 25, 26]. These specific antibodies are synthesised in the brain [24]. The CSF-serum quotient (L₅₀) allows to differentiate between a blood-derived and a pathological brain-derived specific antibody fraction in CSF, taking into account individual changes in the blood/CSF barrier function [24, 25, 26, 27]. Therefore it is necessary to confirm the presence of antibodies against MV using ELISA both in CSF and in the serum. During measles myelitis or encephalitis an intrathecal synthesis of antibodies against MV in CSF takes place. Due to the fact that specific antibodies can pass the blood-cerebrospinal fluid barrier by diffusion from serum to CSF it is necessary to determine the relative CSF/serum quotient (CSQ_{rel}, synonym: antibody specificity index) [24, 25, 26]. The quotient is calculated from the amount of specific anti-measles virus IgG antibodies in total CSF IgG in proportion to the amount of specific IgG antibodies in total serum IgG. During conversion the CSF/serum quotient of the pathogen-specific IgG-antibody concentrations CSQ_{path-spec} (IgG) is put into relation to the CSF/serum quotient of the total IgG concentrations CSQ_{total} (IgG) [27]. A relative CSQ result above 1.5 indicates the production of specific antibodies in the central nervous system (CNS) and the involvement of the CNS in the disease [25, 26].

With respect to the severe complications known from measles infections, the Robert Koch Institute in Germany recommends vaccinating small children, with a first shot between the age of 11 to 14 months and a second between 15 and 23 months [2, 4, 10, 11]. Neutralisation activity and persistence of antibodies are induced in response to the immunisation [6, 15].

Life-long immunity is generally developed. However, antibody levels are 8 to 10 times lower in post-vaccination sera than in convalescent sera [6, 19, 28]. A passive immunisation with specific immunoglobulin concentrates is usually given to immunosuppressed seronegative individuals, such as tumour patients and recipients of transplants, as well as to seronegative pregnant women after exposure to the virus.

The European Regional Office of the WHO aims at eliminating measles from the region in the following years by area-wide vaccination campaigns [4, 9, 10]. This is expected to limit the number of apparent infections and especially of severe courses of the disease. For the diagnosis of the remaining cases of measles infection and of infections acquired outside Europe as well as for the clarification of atypical courses of the disease in partly immunised patients the antibody determination in serum and CSF will be of growing importance [3, 5, 8, 9].



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Avidity determination of antibodies against Measles Viruses (IgG)
Test instruction


ORDER NO.	ANTIBODIES AGAINST	IG CLASS	SUBSTRATE	FORMAT
EI 2610-9601-1 G	Measles viruses	IgG	Ag-coated microplate wells	96 x 01 (96)

Background

The differentiation between fresh and long-standing infections is one of the greatest challenges in serology. Until now this was based mainly on determination of specific antibodies of the immunoglobulin class IgM, which generally only appear initially. However, the detection of these antibodies is often unreliable and problematic due to interfering factors such as persistence of the IgM response, too weak or delayed IgM production, and unspecific IgM production through polyclonal B-cell stimulation.

In recent years additional determination of the antibody avidity has become an established method for identification of primary infections. The immune system reacts to an infection by first forming low-avidity antibodies. With continued disease duration, IgG that are more precisely adapted to the antigens are produced – the avidity increases. If high-avidity IgG are detectable in the serum, it can be assumed that the infection is at a late stage.

Contents of the test system: EI 2610-9601-1 G:

Component	Colour	Format	Symbol
1. Test kit Anti-Measles Viruses ELISA (IgG, order number EI 2610-9601 G)	---	---	---
2. Positive control HA High-avidity anti-Measles (IgG, human), ready for use	red	1 x 1.3 ml	POS CONTROL HA
3. Positive control LA Low-avidity anti-Measles (IgG, human), ready for use	blue	1 x 1.3 ml	POS CONTROL LA
4. Urea solution for Anti-Measles ELISA, ready for use	yellow	1 x 12 ml	UREA
5. Phosphate buffer ready for use	light blue	1 x 12 ml	PBS BUFFER
6. Test instruction	---	1 booklet	---
LOT	Storage temperature		
Lot	Unopened usable until		
In-vitro determination			

Storage and stability: The test kit has to be stored at a temperature between 2°C to +8°C. Do not freeze. Unopened, all test kit components are stable until the indicated expiry date.

Waste disposal: Patient samples, calibrators, controls and incubated microplate strips should be handled as infectious waste. All reagents are to be disposed of according to official regulations.

Preparation and stability of the reagents

Note: All reagents must be brought to room temperature (+18 C to +25 C), approx. 30 minutes before use. After first use, the reagents are stable until the indicated expiry date if stored at +2 C to +8 C and protected from contamination, unless stated otherwise below.

- Controls: Ready for use. The reagents must be mixed thoroughly before use.
- Urea solution: Ready for use. The urea solution included in this test system may only be used for the avidity determination of antibodies against Measles.
- Phosphate buffer: Ready for use.

Warning: The calibrators and controls used have been tested negative for HBsAg, anti-HCV, anti-HIV-1 and anti-HIV-2 using enzyme immunoassays and indirect immunofluorescence methods. Nonetheless, all materials should be treated as being a potential infection hazard and should be handled with care. Some of the reagents contain the toxic agent sodium azide. Avoid skin contact.

Preparation and stability of the patient samples

Sample material: Human serum or EDTA, heparin or citrate plasma.

Stability: Patient samples to be investigated can generally be stored at +2°C to +8 C for up to 14 days. Diluted samples should be incubated within one working day.

Sample dilution: Patient samples are diluted 1:101 sample buffer. For example: dilute 10 µl serum to 1.0 ml sample buffer and mix well by vortexing (sample pipettes are not suitable for mixing).

NOTE: The controls are prediluted and ready for use, do not dilute them.



Incubation

Sample incubation:
(1. step)

Transfer 100 µl of the controls or diluted patient samples into the individual microplate wells according to the pipetting protocol.
Incubate for 30 minutes at room temperature (+18°C to +25°C).

Wash:

Manual: Empty the wells and subsequently wash 1 time using 300 µl of working strength wash buffer.
Automatic: Wash reagent wells 1 time with 450 µl of working strength wash buffer (program setting: e.g. TECAN Columbus Washer "Overflow Modus").

Leave the wash buffer in each well for 30 to 60 seconds per washing cycle, then empty the wells. After washing (manual and automated tests), thoroughly dispose of all liquid from the microplate by tapping it on absorbent paper with the openings facing downwards to remove all residual wash buffer. Free positions on the microplate strip should be filled with blank wells of the same plate format as that of the parameter to be investigated.

Urea incubation:
(2. step)

Pipette 200 µl of urea solution into each of the microplate wells of the first microtiter strip and 200 µl of phosphate buffer into each of the microplate wells of the second microtiter strip.
Incubate for 10 minutes at room temperature (+18°C to +25°C).

Wash:

Empty the wells. Wash as described above, but wash 3 times using working strength wash buffer for each wash.

Attention: Residual liquid (> 10 µl) in the reagent wells after washing can interfere with the substrate and lead to false low extinction values.
Insufficient washing (e.g. less than 3 wash cycles, too small wash buffer volumes, or too short reaction times) can lead to false high extinction values.

Conjugate incubation:
(3. step)

Pipette 100 µl of enzyme conjugate (peroxidase-labelled anti-human IgG) into each of the microplate wells.
Incubate for 30 minutes at room temperature.

Wash:

Empty the wells. Wash as described above, but wash 3 times using working strength wash buffer for each wash.

Substrate incubation:
(4. step)

Pipette 100 µl of chromogen/substrate solution into each of the microplate wells.
Incubate for 15 minutes at room temperature (+18°C to +25°C) protect from direct sunlight.

Stopping the reaction:

Pipette 100 µl of stop solution into each of the microplate wells in the same order and at the same speed as the chromogen/substrate solution was introduced.

Measurement:

Photometric measurement of the colour intensity should be made at a wavelength of 450 nm and a reference wavelength between 620 nm and 650 nm within 30 minutes of adding the stop solution. Prior to measuring, slightly shake the microplate to ensure a homogeneous distribution of the solution.



Pipetting protocol

	1	2	3	4	5	6	7	8	9	10	11	12
A	pos HA	pos HA	P 7	P 7	P 15	P 15						
B	pos LA	pos LA	P 8	P 8	P 16	P 16						
C	P 1	P 1	P 9	P 9	P 17	P 17						
D	P 2	P 2	P 10	P 10	P 18	P 18						
E	P 3	P 3	P 11	P 11								
F	P 4	P 4	P 12	P 12								
G	P 5	P 5	P 13	P 13								
H	P 6	P 6	P 14	P 14								

The above pipetting protocol is an example of the avidity determination of IgG antibodies in 18 patient samples (P 1 to P 18).

Controls (pos HA and pos LA) as well as the patient samples have been incubated in duplicate in one well each of two different microtiter strips. The reagent wells of the microtiter strips 1, 3, 5 etc. are treated with urea solution after the incubation with patients samples, the reagent wells of the microtiter strips 2, 4, 6 etc. are treated with phosphate buffer.

The wells can be broken off individually from the strips. This makes it possible to adjust the number of test substrates used to the number of samples to be examined and minimises reagent wastage.

Both positive controls with high-avidity and low-avidity antibodies serve as internal controls for the reliability of the test procedure. They should be assayed with each test run.

Calculation of results

The presence of low-avidity antibodies in a patient's serum has been proved if the ELISA extinction value is considerably reduced by urea treatment. For an objective interpretation the relative avidity index (RAI) is calculated and expressed in percent using the extinction values with and without urea treatment.

$$\frac{\text{Extinction of the sample with urea treatment} \times 100}{\text{Extinction of the sample without urea treatment}} = \text{relative avidity index (RAI) in \%}$$

The upper limit of the range of low-avidity antibodies (cut-off value) recommended by EUROIMMUN is 40% RAI. Values below the indicated cut-off are to be considered as an indication of low-avidity antibodies, values between 40% and 60% RAI as equivocal, values above 60% RAI as an indication of high-avidity antibodies. If a result is classified as equivocal, it is recommended to collect a second sample not less than 7 days later and to test it together with the first sample.

- RAI < 40%: indication of low-avidity antibodies
- RAI 40% - 60%: equivocal
- RAI > 60%: indication of high-avidity antibodies

Reliable results in the measurement of IgG antibody avidity can only be yielded if the patient sample contains a diagnostically significant concentration of specific antibodies.

Generally, the determination of the relative avidity index is not helpful in samples which have an O.D. of <0.140 after incubation without urea treatment.