

# **SECRETORY IGA ELISA**

**Product Data Sheet** 

Cat. No.: RTC021R

For Research Use Only

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- This kit is manufactured by:
  BioVendor Laboratorní medicína a.s.
- Use only the current version of Product Data Sheet enclosed with the kit!

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### 1. INTENDED USE

A solid-phase enzyme immunoassay for the quantitative determination of secretory IgA in biological fluids. This kit is designed for measurement of secretory IgA in biological fluids. For possibility of use with other sample types, please, refer to Application Notes (on request). The kit contains reagents sufficient for 96 determinations and allows to analyze 41 unknown samples in duplicates.

### 2. STORAGE, EXPIRATION

Store the whole kit at 2 to 8 °C upon receipt until the expiration date. After opening the pouch keep unused microtiter wells TIGHTLY SEALED BY ADHESIVE TAPE (INCLUDED) to minimize exposure to moisture.

### 3. INTRODUCTION

Secretory IgA (sIgA) is the main immunoglobulin present on mucosal surfaces. Ca. 90% of sIgA is produced locally and does not penetrate into blood circulation. sIgA is considerably different from serum IgA, as this complex protein consists of 3 completely different molecules. Two or four molecules of immunoglobulin A with molecular weight 160 kDa are joined by J-chain (16 kDa) and attached to the secretory component (80kDa); the formation of this complex occurs during transepithelial transport of polymeric IgA.

slgA plays a pivotal role in local immunity by blocking bacterial and viral adhesion and invasion through epithelial tissues. Determination of slgA concentration allows to evaluate the local immunity status in stomatology, ophthalmology, respiratory diseases, gastroenterology, gynaecology. The slgA in saliva can be also used as noninvasive mass screening for selective IgA deficiency.

Elevation of slgA in serum is occasionally observed in so autoimmune diseases and several tumours.

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#### 4. TEST PRINCIPLE

This test is based on two-site sandwich enzyme immunoassay principle. Tested specimen is placed into the microwells coated by specific murine monoclonal to human secretory IgA-antibodies. Antigen from the specimen is captured by the antibodies coated onto the microwell surface. Unbound material is removed by washing procedure. Second antibodies - murine monoclonal to human IgA alpha chain, labelled with peroxidase enzyme, are then added into the microwells. After subsequent washing procedure, the remaining enzymatic activity bound to the microwell surface is detected and quantified by addition of chromogen-substrate mixture, stop solution and photometry at 450 nm. Optical density in the microwell is directly related to the quantity of the measured analyte in the specimen.

# Equipment and material required but not provided

- Distilled or deionized water:
- Automatic or semiautomatic multichannel micropipettes, 100-250 μl, is useful but not essential;
- Calibrated micropipettes with variable volume, range volume 25-250 μl;
- Dry thermostat for 37°C +/- 0.1°C
- Calibrated microplate photometer with 450 nm wavelength and OD measuring range 0-3.0.

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### WARNINIGS AND PRECAUTIONS

- This kit is intended for in vitro diagnostic use only.
- INFECTION HAZARD: There is no available test methods that can absolutely assure that
  Hepatitis B and C viruses, HIV-1/2, or other infectious agents are not present in the reagents
  of this kit. All human products, including patient samples, should be considered potentially
  infectious. Handling and disposal should be in accordance with the procedures defined by
  an appropriate national biohazard safety guidelines or regulations.
- Avoid contact with stop solution containing acidic solution. It may cause skin irritation and burns.
- Wear disposable latex gloves when handling specimens and reagents. Microbial contamination of reagents may give false results.
- Do not use the kit beyond the expiration date.
- All indicated volumes have to be performed according to the protocol. Optimal test results are only obtained when using calibrated pipettes and microplate readers.
- Do not smoke, eat, drink or apply cosmetics in areas where specimens or kit reagents are handled.
- Chemicals and prepared or used reagents have to be treated as hazardous waste according to the national biohazard safety guidelines or regulations.
- Do not mix reagents from different lots.
- Replace caps on reagents immediately. Do not swap caps.
- Do not pipette reagents by mouth.
- Specimens must not contain any AZIDE compounds they inhibit activity of peroxidase.
- Safety Data Sheet for this product is available upon request directly from BioVendor Laboratorní medicína a.s.
- The Safety Data Sheet fit the requirements of EU Guideline 91/155 EC.

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# 7. REAGENT SUPPLIED

	Description		Qty	Units	Colour code	Stability of opened/diluted components
1	secretory IgA EIA strips, 8x12 wells	polystyrene microwells coated with murine monoclonal antibodies to human secretory IgA	1	pcs		until exp.date
2	Calibrator set, 1 ml each. The set contains 6 calibrators: 0; 2; 20; 40; 100, 400 µg/ml	human secretory IgA diluted in tris buffered BSA solution, preservative - 0,01% Bronidox L,0,01% 2-Methyl-4 isothiazolin-3-one-hydrochloride; also contains bright blue dye	6	pcs	bright blue (C1 - colourless)	2 months
3	Control serum (1 ml)	dilution of preselected human serum, with high content of secretory IgA with BSA solution; preservative - 0,01% Bronidox L, 0,01% 2-Methyl-4-isothiazolin-3-one-hydrochloride, colourless	1	pcs	colourless	2 months
4	Conjugate, 14 ml	aqueous solution of murine monoclonal to human IgA alfa chain coupled with horseradish peroxidase diluted on phosphate buffered solution with casein from bovine milk and detergent (Tween-20), contains 0,1% phenol as preservative and bright red dye	1	pcs	bright red	until exp.date
5	red EIA buffer 22 ml	phosphate buffered saline with casein from bovine milk and detergent (Tween-20), contains 0,1% phenol as preservative; contains red dye	1	pcs	red	until exp.date
6	EIA buffer 100 ml	phosphate buffered saline with casein from bovine milk and detergent (Tween-20), contains 0,1% phenol as preservative and blue dye	1	pcs	blue	until exp.date
7	Substrate solution, 14 ml	ready-to-use single- component tetramethylbenzidine (TMB) solution.	1	pcs	colourless	until exp.date
8	Washing solution concentrate 26x, 22 ml	aqueous solution of sodium chloride and detergent (Tween 20), contains proClin300 as a preservative	1	pcs	colourless	Concentrate - until exp.date Diluted washing solution - 45 days at 2-8 °C or 15 days at RT
9	Stop solution, 14 ml	5,0% vol/vol solution of sulphuric acid	1	pcs	colourless	until exp.date
10	Plate sealing tape		2	pcs		N/A

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### 8. SPECIMENT COLLECTION AND STORAGE

This kit is intended for use with serum or plasma (ACD- or heparinized), saliva, urine, bronchoalveolar fluid, nasal wash, vagin secret, breast milk. Grossly hemolytic, lipemic, or turbid samples should be avoided.

Specimens may be stored for up to 48 hours at 2-8 °C before testing. Calibrators and control sample(s) - only one freezing/thawing cycle is allowed.

### 9. TEST PROCEDURE

### 9.1 Preparation of reagent

- All reagents (including unsealed microstrips) should be allowed to reach room temperature (+18 to +25°C) before use.
- All reagents should be mixed by gentle inversion or vortexing prior to use. Avoid foam formation.
- It is recommended to spin down shortly the tubes with calibrators on low speed centrifuge.
- Prepare washing solution from the concentrate WASH SOLN 26X by 26 dilutions in distilled water.

#### 9.2 Procedural Note

It is recommended that pipetting of all calibrators and samples should be completed within 3 minutes.

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# 9.3 Assay procedure

1	Put the desired number of microstrips into the frame; allocate 14 wells for the calibrators CAL 1 - 6 and control samples CONTROL and two wells for each unknown sample. DO NOT REMOVE ADHESIVE SEALING TAPE FROM UNUSED STRIPS. NOTE: the calibrator/control and unknown sample wells are filled differently.		
2	Dilute samples using buffer SAM DIL (EIA buffer) 101 fold. See table M for dilution modes and factors for different types of analyzed material. Do not dilute control sample and calibrators.		
3	If suggested analyte concentration in the sample exceeds the highest calibrator, additionally dilute this sample accordingly, using SAM DIL (EIA buffer). Use of other buffers or reagents for sample dilution may lead to incorrect measurement.		
4	Pipet 190 µI of red EIA buffer into the wells allocated for saliva. For other tested materials, see table M for the volume of red EIA buffer.		
5	Pipet 100 µl of calibrators CAL 1 - 6 and control samples CONTROL into allocated wells. For testing of saliva pipet 10 µl of the unknown sample into the allocated wells. See table M for the volumes of other materials. Carefully mix the contents of the wells by short horizontal rotating of the plate for 5-7 seconds and cover the wells by plate adhesive tape (included into the kit).		
6	Incubate 90 minutes at 37 °C .		
7	Prepare washing solution by 26x dilution of washing solution concentrate WASH SOLN 26X by distilled water. Minimal quantity of washing solution should be 250 µl per well. Wash strips 3 times		
8	Dispense 100 µl of ENZ CONJ into the wells. Cover the wells by plate adhesive tape.		
9	Incubate 30 minutes at 37 °C .		
10	Wash the strips 5 times.		
11	Dispense 100 µl of SUB TMB into the wells		
12	Incubate 10-20 minutes at 18-25 °C		
13	Dispense 100 µl of STOP SOLN into the wells.		
14 15	Measure OD (optical density) at 450 nm.		
16	Set photometer blank on first calibrator  Apply point-by-point method for data reduction. Use Calculation factor listed in table M to calculate analyte concentration in different material types.		

# 9.4 Handing notes

Calibrators and control sample(s) - only one freezing/thawing cycle is allowed

# 9.5 Sample Processing

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Material type	Notes on material collection, storage and handling	Sample dilution example	red EIA buffer into the well, µl	Sample into the well, µl	Calculation factor
blood serum or plasma	Grossly hemolytic, lipemic, or turbid samples should be avoided and should be treated by centrifugation before testing.	5 µl of sample + 500 µl of diluent	0	100	0.05
saliva	Do not eat and avoid vigorous physical activity 60 minutes before sample collection. Document consumption of alcohol, caffeine, nicotine, and prescription/over-the-counter medications within the prior 12 hours. Document the presence of oral diseases or injury. Before collecting saliva, rinse mouth with water to remove food residue and wait at least 10 minutes after rinsing to avoid sample dilution. Before assaying, saliva samples should be frozen at -20 °C overnight, then thawed and centrifuged at 1000 g for ten minutes to remove particulate matter. Undiluted specimens may be stored refrigerated at (2-8 °C) for 5 days. If storage time exceeds 5 days, store frozen at (-20 °C) for up to one month. Avoid repeated freeze thaw cycles.	5 µl of sample + 500 µl of diluent	190	10	1.0
urine	Use a sterile container to collect urine samples. Remove any particulates by centrifugation for 15 minutes at 1000xg, 2-8 °C and assay immediately or aliquot and store samples at -20°C or -80 °C for up to one month. Avoid repeated freeze-thaw cycles. Centrifuge again before assaying to remove any additional precipitates that may appear after storage.	10 µl of sample + 500 µl of diluent	0	100	0.025
bronchoalveolar fluid	Turbid samples should give incorrect measurement results and should be treated by centrifugation before testing.	5 μl of sample + 500 μl of diluent	0	100	0.05
nasal wash	Turbid samples should give incorrect measurement results and should be treated by centrifugation before testing.	5 µl of sample + 500 µl of diluent	80	20	0.25
vaginal secret	Turbid samples should give incorrect measurement results and should be treated by centrifugation before testing.	5 µl of sample + 500 µl of diluent	90	10	0.5
breast milk	Turbid samples should give incorrect measurement results and should be treated by centrifugation before testing.	5 µl of sample + 2500 µl of diluent	195	5	10

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#### QUALITY CONTROL

It is recommended to use control samples according to state and federal regulations. The use of control samples is advised to assure the day to day validity of results.

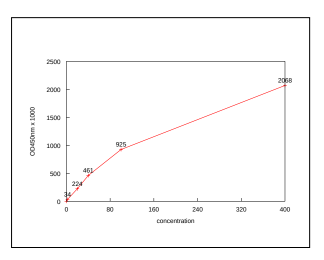
The test must be performed exactly as per the manufacturer's instructions for use. Moreover the user must strictly adhere to the rules of GLP (Good Laboratory Practice) or other applicable federal, state, and local standards and/or laws. This is especially relevant for the use of control reagents. It is important to always include, within the test procedure, a sufficient number of controls for validating the accuracy and precision of the test.

The test results are valid only if all controls are within the specified ranges and if all other test parameters are also within the given assay specifications.

### 11. CALCULATION

- Calculate the mean absorbance values (OD450) for each pair of calibrators and samples.
- Plot a calibration curve on graph paper: OD versus secretory IgA concentration.
- Determine the corresponding concentration of secretory IgA in unknown samples from the calibration curve. Manual or computerized data reduction is applicable on this stage. Point-by-point or linear data reduction is recommended due to non-linear shape of curve.
- Below is presented a typical example of a standard curve with the BioVendor Assay. Not for calculations!

Calibrators	Value	Absorbance Units (450
		nm)
CAL 1	0 μg/ml	0.10
CAL 2	2 μg/ml	0.14
CAL 3	20 µg/ml	0.33
CAL 4	40 µg/ml	0.57
CAL 5	100	1.03
CAL 5	μg/ml	1.00
CAL 6	400	2.17
CAL 0	μg/ml	2.11



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### 12. EXPECTED VALUES

Therapeutical consequences should not be based on results of IVD methods alone – all available clinical and laboratory findings should be used by a physician to elaborate therapeutically measures. Each laboratory should establish its own normal range for secretory IgA. Based on data obtained by BioVendor, the following normal range is recommended (see below). NOTE: the patients that have received murine monoclonal antibodies for radioimaging or immunotherapy develop high titered anti-mouse antibodies (HAMA). The presence of these antibodies may cause false results in the present assay. Sera from HAMA positive patients should be treated with depleting adsorbents before assaying

	Units			
Sex, age	μg/ml			
	Lower limit	Upper limit		
serum	1.6	5.0		
saliva	57	260		
urine	0.5	2.7		
breast milk	800	-		

# 13. PERFORMANCE CHARACTERISTICS

### 13.1 Sensitivity

Sensitivity of the assay was assessed as being 0.6 µg/ml.

### 13.2 Cross-reactivity

Analyte	Cross-reactivity, % wt/wt
IgG	<0.1
IgM	<0.1
IgE	<0.1

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### 13.3 Linearity

Linearity was checked by assaying dilution series of 5 samples with different secretory IgA concentrations. Linearity percentages obtained ranged within 90 to 110%.

# 13.4 Recovery

Recovery was estimated by assaying 5 mixed samples with known secretory IgA concentrations. The recovery percentages ranged from 90 to 110%.

### 14. REFERENCES

- 1. Heiddis B. Valdimarsdottir and Arthur A. Stone Psychosocial Factors and Secretory Immunoglobulin A. Critical Reviews in Oral Biology & Medicine, Jan 1997; 8: 461 474.
- 2. Amir H Abdul Latiff and Michael A Kerr The clinical significance of immunoglobulin A deficiency. Ann Clin Biochem, Mar 2007; 44: 131 139.

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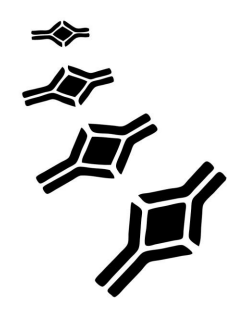
# 15. EXPLANATION OF SYMBOLS

Symbol	English
<b>(</b> €	European Conformity
	Consult instructions for use
IVD	In vitro diagnostic device
REF	Catalogue number
LOT	Lot. No. / Batch code
RUO	For research use only
Σ	Contains sufficient for <n> tests/</n>
1	Storage Temperature
2	Expiration Date
***	Legal Manufacturer
Distributed by	Distributor

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