

#### Manual

# *IDK®* slgA ELISA

For the determination of secretory IgA in saliva and stool

Valid from 2020-11-03



KR8870









KR8870.20



 $\frac{2}{20 \times 96}$ 



Immundiagnostik AG, Stubenwald-Allee 8a, 64625 Bensheim, Germany

Fax: + 49 6251 70190-363

e.mail: info@immundiagnostik.com www.immundiagnostik.com

## **Table of Contents**

1.	INTENDED USE	2
2.	INTRODUCTION	2
3.	MATERIAL SUPPLIED	2
4.	MATERIAL REQUIRED BUT NOT SUPPLIED	3
5.	STORAGE AND PREPARATION OF REAGENTS	3
6.	STORAGE AND PREPARATION OF SAMPLES	4
	Sample stability	4
	Saliva	
	Extraction of the stool samples	5
	Dilution of stool samples	6
7.	ASSAY PROCEDURE	6
	Principle of the test	6
	Test procedure	7
8.	RESULTS	8
9.	LIMITATIONS	9
10.	QUALITY CONTROL	9
	Reference range	9
11.	PERFORMANCE CHARACTERISTICS	9
	Accuracy – Precision	9
	Analytical sensitivity	
12.	PRECAUTIONS	10
13.	TECHNICAL HINTS	10
14.	GENERAL NOTES ON THE TEST AND TEST PROCEDURE	11
15.	REFERENCES	11
	General literature	11
	Literature using K8870	11

#### 1. INTENDED USE

This Immundiagnostik AG assay is an enzyme immunoassay intended for the quantitative determination of secretory IgA (sIgA) in saliva and stool.

For research use only. Not for use in diagnostic procedures.

#### 2. INTRODUCTION

Secretory IgA (sIgA) consists of two IgA monomers joined by the J-chain and an additional secretory component. It is secreted in plasma cells located in the *lamina propia* of mucosal membranes. Synthesis of sIgA is independent from the synthesis of serum IgA. This means that lack of serum IgA does not necessarily correlate with a lack of sIgA1. Secretory IgA is the major immunoglobulin in saliva, tears, colostrum, nasal mucous, mother's milk, tracheobronchial and gastrointestinal secretes. It plays a major role in preventing adherence of microorganisms to mucosal sites, in activation of the alternative complement pathway and in activating inflammatory reactions.

### 3. MATERIAL SUPPLIED

Aut no	Label Kit components	Quantity for cat. no.		
Art. no.		Kit components	KR8870	KR8870.20
KR8870	PLATE	Microtiter plate, pre-coated	12 x 8 wells	20 x 12 x 8 wells
KR0001.C.100	WASHBUF	Wash buffer concentrate, 10x	2 x 100 ml	-
KR8870	CONJ	Conjugate concentrate, peroxidase-labelled (mouse anti-slgA)	1 x 200 μl	15 x 200 μl
KR8870	STD	Standards, lyophilised (0; 22.2; 66.6; 200; 600 ng/ml)	2 x 5 vials	25 x 5 vials
KR8870	CTRL1	Control, lyophilised (see specification for range)	2 x 1 vial	25 x 1 vial
KR8870	CTRL2	Control, lyophilised (see specification for range)	2 x 1 vial	25 x 1 vial

Aut no	Label	Vit components	Quantity for cat. no.	
Art. no. L	Labei	Kit components	KR8870	KR8870.20
KR0002.15	SUB	Substrate (tetramethyl- benzidine), ready-to-use	1 x 15 ml	20 x 15 ml
KR0003.15	STOP	Stop solution, ready-to-use	1 x 15 ml	20 x 15 ml
KR6999.C.100	IDK Extract®	Extraction buffer concentrate IDK Extract® 2.5x	2 x 100 ml	_

For reorders of single components, use the catalogue number followed by the label as product number.

## 4. MATERIAL REQUIRED BUT NOT SUPPLIED

- Ultrapure water\*
- Stool sample application system such as cat. no.: KR6998SAS
- Calibrated precision pipettors and 10–1000 µl single-use tips
- · Foil to cover the microtiter plate
- · Horizontal microtiter plate shaker
- Multi-channel pipets or repeater pipets
- Centrifuge, 3000 g
- Vortex
- Standard single-use laboratory glass or plastic vials, cups, etc.
- Microtiter plate reader (required filters see chapter 7)
  - \* Immundiagnostik AG recommends the use of ultrapure water (water type 1; ISO 3696), which is free of undissolved and colloidal ions and organic molecules (free of particles > 0.2  $\mu$ m) with an electrical conductivity of 0.055  $\mu$ S/cm at 25 °C ( $\geq$  18.2 M $\Omega$ cm).

## 5. STORAGE AND PREPARATION OF REAGENTS

- To run the assay more than once, ensure that reagents are stored at the conditions stated on the label. Prepare only the appropriate amount necessary for each run. The kit can be used up to 4 times within the expiry date stated on the label.
- Reagents with a volume less than 100 μl should be centrifuged before use to avoid loss of volume.
- Preparation of the wash buffer: The wash buffer concentrate (WASHBUF) has to be diluted with ultrapure water 1:10 before use (100 ml WASHBUF + 900 ml ultrapure water), mix well. Crystals could occur due to high salt con-

centration in the concentrate. Before dilution, the crystals have to be redissolved at room temperature or in a water bath at 37 °C. The **WASHBUF** can be used until the expiry date stated on the label when stored at **2–8 °C. Wash buffer** (1:10 diluted WASHBUF) can be stored in a closed flask at **2–8 °C for 1 month**.

- Preparation of the extraction buffer: The extraction buffer concentrate *IDK Extract*® has to be diluted with ultrapure water 1:2.5 before use (100 ml *IDK Extract*® + 150 ml ultrapure water), mix well. Crystals could occur due to high salt concentration in the concentrate. Before dilution, the crystals have to be redissolved at 37 °C in a water bath. The *IDK Extract*® can be used until the expiry date stated on the label when stored at 2–8 °C. Extraction buffer (1:2.5 diluted *IDK Extract*®) can be stored in a closed flask at 2–8 °C for 4 months.
- The lyophilised standards (STD) and controls (CTRL) can be used at 2–8 °C until the expiry date stated on the label. Before use, the STD and CTRL have to be reconstituted with 500 µl of ultrapure water and mixed by gentle inversion to ensure complete reconstitution. Allow the vial content to dissolve for 10 minutes and then mix thoroughly. Standards and controls (reconstituted STD and CTRL) can be stored at -20 °C for 4 weeks and can be subjected to a maximum of two freeze-thaw cycles.
- Preparation of the conjugate: Before use, the conjugate concentrate (CONJ) has to be diluted 1:101 in wash buffer (100 µl CONJ + 10 ml wash buffer). The CONJ can be used at 2–8 °C until the expiry date stated on the label. Conjugate (1:101 diluted CONJ) is not stable and cannot be stored.
- All other test reagents are ready-to-use. Test reagents can be used until the expiry date (see label) when stored at 2–8°C.

#### 6. STORAGE AND PREPARATION OF SAMPLES

Sample stability

The **sample stability** is as follows:

**Raw stool:** 2 days at room temperature (15–30 °C), 2 days at 2–8 °C or 8 weeks at -20 °C

**Stool extracts (1:100):** 1 day at room temperature (15–30 °C), 7 days at 2–8 °C or 7 days at -20 °C, maximum 2 freeze-thaw cycles

Saliva: 1 day at 2–8 °C, 4 weeks at -20 °C

#### Saliva

To avoid variation in slgA content, take saliva samples always at the same time of the day. No food or liquid should be consumed 30 min before sample collection. Collect saliva samples using salivettes and centrifuge at 3000 *q* for 10 min.

For analysis, the saliva supernatant is diluted 1:2000 in wash buffer, e.g.

10  $\mu$ l saliva supernatant + 990  $\mu$ l wash buffer = dilution I (1:100)

**50 μl** dilution  $I + 950 \mu I$  wash buffer = dilution II (1:20)

Final dilution: 1:2000

For analysis, pipet 100 µl of dilution II per well.

## Extraction of the stool samples

**Extraction buffer** (1:2.5 diluted *IDK Extract*\*) is used as a **sample extraction buffer**. We recommend the following sample preparation:

#### Stool Sample Application System (SAS) (Cat. No.: KR6998SAS)

#### Stool sample tube – Instructions for use

Please note that the dilution factor of the final stool suspension depends on the amount of stool sample used and the volume of the buffer.

## SAS with 1.5 ml sample extraction buffer:

Applied amount of stool: 15 mg
Buffer Volume: 1.5 ml
Dilution Factor: 1:100

Please follow the instructions for the preparation of stool samples using the SAS as follows:

- a) The raw stool sample has to be thawed. For particularly heterogeneous samples we recommend a mechanical homogenisation using an applicator, inoculation loop or similar device.
- b) Fill the **empty stool sample tube** with 1.5 ml **sample extraction buffer** (1:2.5 diluted *IDK Extract*\*) before using it with the sample. **Important:** Allow the sample extraction buffer to reach room temperature.
- c) Unscrew the tube (yellow part of cap) to open. Insert the yellow dipstick into the sample. The lower part of the dipstick has notches which need to be covered completely with stool after inserting it into the sample. Place dipstick back into the tube. When putting the stick back into the tube, excess material

will be stripped off, leaving 15 mg of sample to be diluted. Screw tightly to close the tube.

- d) Vortex the tube well until no stool sample remains in the notches. **Important:** Please make sure that you have a maximally homogenous suspension after shaking. Especially with more solid samples, soaking the sample in the tube with sample extraction buffer for ~10 minutes improves the result.
- e) Allow sample to stand for ~10 minutes until sediment has settled. Floating material like shells of grains can be neglected.
- f) Carefully unscrew the complete cap of the tube including the blue ring plus the dipstick. Discard cap and dipstick. Make sure that the sediment will not be dispersed again.

Dilution I: 1:100

## Dilution of stool samples

The supernatant of the sample preparation procedure (dilution I) is further diluted **1:125 in wash buffer**. For example:

**40 μl** dilution I + 960 μl wash buffer (mix well) = **dilution II** (1:25)

 $200\,\mu l$  dilution II +  $800\,\mu l$  wash buffer (mix well) = dilution III (1:5)

Final dilution: 1:12 500

For analysis, pipet 100 µl of dilution III per well.

## 7. ASSAY PROCEDURE

## Principle of the test

This ELISA is intended for the quantitative determination of secretory IgA in stool and saliva. In a first incubation step, the sIgA in the samples is bound to polyclonal antibodies (rabbit anti human IgA), which are immobilised to the surface of the microtiter wells. To remove all unbound substances, a washing step is carried out. In a second incubation step, a peroxidase-labelled conjugate (mouse anti-sIgA) is added which recognises specifically the bound secretory IgA. After another washing step, to remove all unbound substances, the solid phase is incubated with the substrate, tetramethylbenzidine (TMB). An acidic stop solution is then added to stop the reaction. The colour converts from blue to yellow. The intensity of the yellow colour is directly proportional to the concentration of secretory IgA. A dose response curve of the absorbance unit (optical density, OD) vs. concentration is generated, using the results obtained from the standards. Secretory IgA, present in the samples, is determined directly from this curve.

## Test procedure

Bring all reagents and samples to room temperature (15–30 °C) and mix well.

Mark the positions of standards/controls/samples on a protocol sheet.

Take as many microtiter strips as needed from kit. Store unused strips together with the desiccant bag in the closed aluminium packaging at  $2-8^{\circ}$  C. Strips are stable until expiry date stated on the label.

For automated ELISA processors, the given protocol may need to be adjusted according to the specific features of the respective automated platform. For further details please contact your supplier or Immundiagnostik AG.

We recommend to carry out the tests in duplicate.

1.	Before use, wash the wells <b>5 times</b> with <b>250 µl wash buffer</b> . After the final washing step, the inverted microtiter plate should be firmly tapped on absorbent paper.	
2.	Add each $100\mu l$ standards/controls/diluted samples into the respective wells.	
3.	Cover the strips and incubate for 1 hour on a horizontal shaker* at room temperature (15–30 $^{\circ}$ C).	
4.	Discard the content of each well and wash <b>5 times</b> with <b>250 µl wash buffer</b> . After the final washing step, the inverted microtiter plate should be firmly tapped on absorbent paper.	
5.	Add <b>100 μl conjugate</b> (diluted CONJ) in each well.	
6.	Cover the strips and incubate for <b>1 hour on a horizontal shaker*</b> at room temperature (15–30 $^{\circ}$ C).	
7.	Discard the content of each well and wash 5 times with 250 µl wash buffer. After the final washing step, the inverted microtiter plate should be firmly tapped on absorbent paper.	
8.	Add 100 µl substrate (SUB) in each well.	
9.	Incubate for <b>10–20 minutes**</b> at room temperature (15–30°C) <b>in the dark.</b>	
10.	Add <b>100 μl stop solution</b> (STOP) and mix well.	

Determine **absorption immediately** with an ELISA reader at **450 nm** against 620 nm (or 690 nm) as a reference. If no reference wavelength is available, read only at 450 nm. If the extinction of the highest standard exceeds the range of the photometer, absorption must be measured immediately at **405 nm** against 620 nm as a reference.

#### 8. RESULTS

The following algorithms can be used alternatively to calculate the results. We recommend using the "4 parameter algorithm".

#### 1. 4 parameter algorithm

It is recommended to use a linear ordinate for the optical density and a logarithmic abscissa for the concentration. When using a logarithmic abscissa, the zero standard must be specified with a value less than 1 (e.g. 0.001).

## 2. Point-to-point calculation

We recommend a linear ordinate for the optical density and a linear abscissa for the concentration.

## 3. Spline algorithm

We recommend a linear ordinate for the optical density and a linear abscissa for the concentration.

The plausibility of the duplicate values should be examined before the automatic evaluation of the results. If this option is not available with the programme used, the duplicate values should be evaluated manually.

#### Saliva

The obtained results have to be multiplied by the **dilution factor of 2000** to get the actual concentrations.

#### Stool

The obtained results have to be multiplied by the **dilution factor of 12 500** to get the actual concentrations.

In case **another dilution factor** has been used, multiply the obtained result by the dilution factor used

<sup>\*</sup> We recommend shaking the strips at 550 rpm with an orbit of 2 mm.

<sup>\*\*</sup> The intensity of the colour change is temperature sensitive. We recommend observing the colour change and stopping the reaction upon good differentiation.

#### 9. LIMITATIONS

Samples with concentrations above the measurement range (see definition below) can be further diluted and re-assayed. Please consider this higher dilution when calculating the results.

Samples with concentrations lower than the measurement range (see definition below) cannot be clearly quantified.

The upper limit of the measurement range can be calculated as:

highest concentration of the standard curve × sample dilution factor to be used

The lower limit of the measurement range can be calculated as:

 $LoB \times sample dilution factor to be used$ 

LoB see chapter "Performance Characteristics".

#### **10. QUALITY CONTROL**

Immundiagnostik AG recommends the use of external controls for internal quality control, if possible.

Control samples should be analysed with each run. Results, generated from the analysis of control samples, should be evaluated for acceptability using appropriate statistical methods. The results for the samples may not be valid if within the same assay one or more values of the quality control sample are outside the acceptable limits.

## Reference range

We recommend each laboratory to establish its own reference range.

## 11. PERFORMANCE CHARACTERISTICS

## Accuracy - Precision

## Repeatability (Intra-Assay); n = 20

The repeatability was assessed with 2 stool samples under **constant** parameters (same operator, instrument, day and kit lot).

Sample	Mean value [μg/ml]	CV [%]
1	971.0	5.6
2	1136.0	5.8

#### Reproducibility (Inter-Assay); n = 12

The reproducibility was assessed with 2 stool samples under **varying** parameters (different operators, instruments, days and kit lots).

Sample	Mean value [μg/ml]	CV [%]
1	1 279.6	8.2
2	1 277.4	7.7

## Analytical sensitivity

The following values have been estimated based on the concentrations of the standards without considering possibly used sample dilution factors.

Limit of blank, LoB

2.088 ng/ml

#### 12. PRECAUTIONS

- All reagents in the kit package are for research use only.
- Human materials used in kit components were tested and found to be negative for HIV, Hepatitis B and Hepatitis C. However, for safety reasons, all kit components should be treated as potentially infectious.
- Kit reagents contain sodium azide or ProClin as bactericides. Sodium azide and ProClin are toxic. Substrates for the enzymatic colour reactions are toxic and carcinogenic. Avoid contact with skin or mucous membranes.
- The stop solution consists of diluted sulphuric acid, a strong acid. Although
  diluted, it still must be handled with care. It can cause burns and should be
  handled with gloves, eye protection, and appropriate protective clothing. Any
  spill should be wiped up immediately with copious quantities of water. Do not
  breath vapour and avoid inhalation.

## 13. TECHNICAL HINTS

- Do not interchange different lot numbers of any kit component within the same assay. Furthermore we recommend not assembling wells of different microtiter plates for analysis, even if they are of the same batch.
- Control samples should be analysed with each run.
- Reagents should not be used beyond the expiration date stated on kit label.
- Substrate solution should remain colourless until use.

• To ensure accurate results, proper adhesion of plate sealers during incubation steps is necessary.

- · Avoid foaming when mixing reagents.
- Do not mix plugs and caps from different reagents.
- The assay should always be performed according to the enclosed manual.

#### 14. GENERAL NOTES ON THE TEST AND TEST PROCEDURE

- The guidelines for laboratories should be followed.
- IDK® and IDK Extract® are trademarks of Immundiagnostik AG.
- Incubation time, incubation temperature and pipetting volumes of the components are defined by the producer. Any variation of the test procedure, which is not coordinated with the producer, may influence the results of the test. Immundiagnostik AG can therefore not be held responsible for any damage resulting from incorrect use.
- Warranty claims and complaints regarding deficiencies must be logged within 14 days after receipt of the product. The product should be send to Immundiagnostik AG along with a written complaint.

#### 15. REFERENCES

#### General literature

- 1. Brandtzaeg, P., 2010. Update on mucosal immunoglobulin A in gastrointestinal disease. *Current opinion in gastroenterology*, **26**(6), pp.554–63.
- 2. Corthésy, B., 2012. Autoimmunity Reviews Role of secretory IgA in infection and maintenance of homeostasis. *Autoimmunity Reviews*.

## Literature using K8870

- 3. Kalach, N. et al., 2013. Intestinal permeability and fecal eosinophil-derived neurotoxin are the best diagnosis tools for digestive non-lgE-mediated cow's milk allergy in toddlers. *Clinical chemistry and laboratory medicine: CCLM / FESCC*, **51**(2), pp.351–61.
- 4. Kaur, R. et al., 2012. Antibody in middle ear fluid of children originates predominantly from sera and nasopharyngeal secretions. *Clinical and vaccine immunology: CVI*, **19**(10), pp.1593–6.

 Kabeerdoss, J. et al., 2011. Effect of yoghurt containing Bifidobacterium lactis Bb12® on faecal excretion of secretory immunoglobulin A and human beta-defensin 2 in healthy adult volunteers. *Nutrition journal*, 10(1), p.138.

- 6. Senol, A. et al., 2011. Effect of probiotics on aspirin-induced gastric mucosal lesions. *The Turkish journal of gastroenterology: the official journal of Turkish Society of Gastroenterology*, **22**(1), pp.18–26.
- 7. Chalkias, A. et al., 2011. Patients with colorectal cancer are characterized by increased concentration of fecal hb-hp complex, myeloperoxidase, and secretory IgA. *American journal of clinical oncology*, **34**(6), pp.561–6.
- 8. Mohan, R. et al., 2008. Effects of Bifidobacterium lactis Bb12 supplementation on body weight, fecal pH, acetate, lactate, calprotectin, and IgA in preterm infants. *Pediatric research*, **64**(4), pp.418–22.
- Hofman, L. & Le, T., 2002. Preliminary pediatric reference range for secretory IgA in saliva using an enzyme immunoassay. *Clinical Chemistry*, 48(6, Supplement), p.A169-70.

#### **Used symbols:**

