

# OXA-23 K-SeT



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## ***In vitro* rapid diagnostic test for the detection of OXA-23 carbapenemase in bacterial culture**

**FOR IN VITRO DIAGNOSTIC USE  
FOR PROFESSIONAL USE ONLY**

EN

References: K-15R7, 20 cassettes, buffer, 20 tubes and droppers

### I. INTRODUCTION

*Acinetobacter baumannii* is an important opportunistic and multidrug-resistant Gram-negative bacteria responsible for nosocomial infections in health facilities. If left untreated, this infection can lead to septicemia and death. The carbapenem-hydrolysing oxacillinases (OXAs) are the most commonly reported carbapenem-resistance determinants in *Acinetobacter* spp., particularly in *A. baumannii*. Among the OXAs, OXA-23 is the most prevalent carbapenem-resistance determinant in *A. baumannii* isolates.

OXA-23 has been detected in other bacterial species as chromosomal (*P. mirabilis*, Bonnet et al 2002 and Osterblad et al 2016; *A. radioresistans*) or plasmidic gene (*E. coli*, La et al, 2014), which can constitute reservoirs for horizontal transmission of this resistance factor (Poirel et al 2016). The detection of this resistance factor OXA-23, not only in resistant species but also in carrier species, is therefore of paramount importance in the control of antibiotic resistance in the hospital.

Nowadays, definitive confirmation of OXA-23 relies on molecular amplification analysis and DNA sequencing. These tests are expensive and can only be performed in dedicated environment and by skilled staff, hence limiting their more generalized usage.

The development of new rapid diagnostic tests to track antimicrobial resistance patterns is considered as one of the priority core action by international experts and health authorities.

The OXA-23 K-SeT test aimed at a rapid identification of the OXA-23 carbapenemase (and variants of the OXA-23 group) ensures effective treatment of patients and prevention of spread of OXA-23 *Acinetobacter* spp. carrier, especially in hospitals.

### II. PRINCIPLE OF THE TEST

This test is ready to use and is based on a membrane technology with colloidal gold nanoparticles. A nitrocellulose membrane is sensitized with a monoclonal antibody directed against one epitope of the OXA-23 carbapenemase. Another monoclonal antibody directed against a second epitope of the OXA-23 carbapenemase is conjugated to colloidal gold particles. This conjugate is dried on a membrane.

This test is aimed at the detection of OXA-23 like carbapenemases in a single bacterial colony growing on agar plate. The sample must be diluted in the dilution buffer supplied with the test. When the provided buffer containing the resuspended bacteria comes into contact with the strip, the solubilized conjugate migrates with the sample by passive diffusion and both the conjugate and sample material come into contact with the anti-OXA-23 antibody that it is adsorbed onto the nitrocellulose strip. If the sample contains the OXA-23 carbapenemase, the conjugate-OXA-23 complex will remain bound to the anti-OXA-23 antibody adsorbed onto the nitrocellulose and a red line will develop. Solution continues to migrate to reach a second reagent (control reagent) that binds the migration control conjugate, thereby producing a red control line that confirms that the test is valid. Result is visible within 15 minutes.

### III. REAGENTS AND MATERIALS

#### 1. OXA-23 K-SeT (20)

20 sealed pouches containing one device and one desiccant. Each device contains one sensitized strip.

#### 2. LY-A buffer vial (15 mL)

Saline solution buffered to pH 7.5 containing TRIS, Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub> (<0,1%) and a detergent.

#### 3. Instruction for use (1)

#### 4. Semi-rigid disposable collection tubes with droppers (20)

### IV. SPECIAL PRECAUTIONS

- All operations linked to the use of the test must be performed in accordance with Good Laboratory Practices (GLP).
- All reagents are for *in vitro* diagnostic use only.
- Pouch must be opened with care.
- Avoid touching nitrocellulose with your fingers.
- Wear gloves when handling samples.

- Never use reagents from another kit.
- Green lines indicate immunoreagents adsorption sites. Green colour disappears during the test.
- Reagents' quality cannot be guaranteed beyond their shelf-life dates or if reagents are not stored under required conditions as indicated in the insert.

### V. WASTE DISPOSAL

- Dispose of gloves, swabs, test tubes and used devices in accordance with GLP.
- Each user is responsible for the management of any waste produced, and must ensure that it is disposed of in accordance with the applicable legislation.

### VI. STORAGE

- An unopened pouch may be kept at between 4 and 30°C and used until the shelf-life date indicated on the packaging. Once the pouch is opened, run the test immediately.
- Avoid freezing devices and buffer.

### VII. SPECIMEN HANDLING AND COLLECTION

Specimens to be tested should be obtained and handled by standard microbiological methods.

Make sure that the specimens are not treated with solutions containing formaldehyde or its derivatives.

### VIII. PROCEDURE

#### PREPARATIONS OF THE TEST:

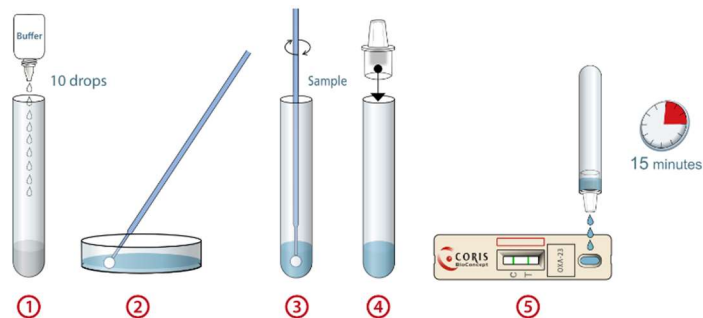
Allow kit components, in unopened packaging, and specimens (in case the plate containing colony to be tested was kept at 4°C) to reach room temperature (15-30°C) before performing a test.

Open the pouch and remove the device. Once opened, run the test immediately. Indicate the patient's name or specimen number on the device (one device per sample).

#### SPECIMEN PREPARATION PROCEDURE:

We recommend the use of fresh bacterial colonies for optimal test performance.

1. Prepare one semi-rigid tube provided in the kit and add 10 drops of LY-A buffer in the tube.
2. Harvest bacteria by taking one colony with a disposable bacteriological loop and dip the loop in the bottom of the semi-rigid tube containing the buffer.
3. Stir thoroughly to homogenize the solution.
4. Insert tightly the dropper on the semi-rigid tube.
5. Invert the test tube and add slowly 3 drops of diluted sample into the sample well of the cassette.
6. Allow to react for 15 min maximum and read the result.



Positive results may be reported as soon as the test and control lines become visible. **Do not take the appearance of new lines into account after the reaction time is passed.**

**The result must be read on still wet strip.**

### IX. INTERPRETING RESULTS

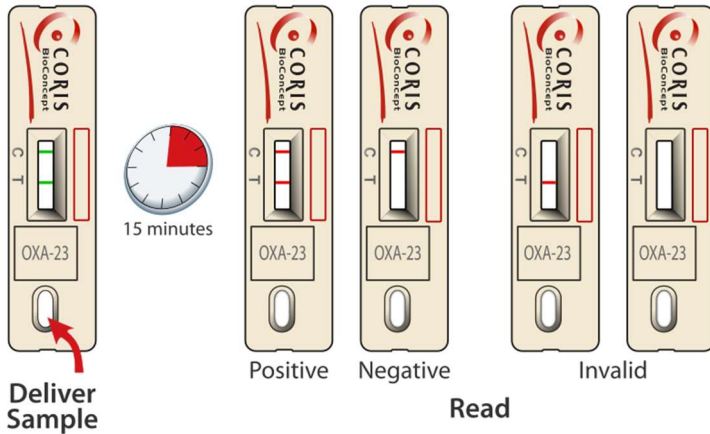
The results are to be interpreted as follows:

**Negative test result:** a reddish-purple line appears across the central reading window at the Control line (C) position. No other band is present.

**Positive test result:** in addition to a reddish-purple band at the Control line (C), a visible reddish-purple band appears at the Test line position (T). Intensity of the test line may vary according to the quantity of antigens present in the sample. Any reddish-purple line (T), even weak, should be considered as a positive result.

**Invalid test result:** The absence of a Control line indicates a failure in the test procedure. Repeat invalid tests with a new test device.

Note: during the drying process, a very faint shadow may appear at the Test line position. It should not be regarded as a positive result.



## X. PERFORMANCE

### A. Detection Limit

The detection limit was determined with a purified recombinant OXA-23 protein and has been evaluated at 0,156 ng/mL.

### B. Validation on collection of reference strains

The OXA-23 K-SeT was evaluated on a collection of 108 clinical isolates of carbapenem-resistant *Acinetobacter* spp. fully characterized resistance mechanisms to beta-lactams by phenotypic and molecular tests (Germany).

108 strains	35 strains tested positive with the OXA-23 K-SeT	35 strains carrying OXA-23 carbapenemase	<i>Acinetobacter baumannii</i> , <i>Acinetobacter pittii</i> , <i>Acinetobacter nosocomialis</i> , <i>Acinetobacter radioresistens</i>
	73 strains tested negative with the OXA-23 K-SeT	68 strains carrying a non-OXA-23 carbapenemase	OXA-40, OXA-51, OXA-58, OXA-143, OXA-235
		5 strains carrying class B carbapenemases	Including VIM-2, NDM-1, NDM-2

### C. Repeatability and reproducibility

To check intra-batch accuracy (repeatability), the same positive samples and a buffer solution were processed 15 times on kits of the same production batch in the same experimental conditions. All observed results were confirmed as expected.

To check inter-batch accuracy (reproducibility), some samples (positive and buffer) were processed on kits from three different production batches. All results were confirmed as expected.

## XI. LIMITS OF THE KIT

The test is qualitative and cannot predict the quantity of antigens present in the sample. Clinical presentation and other test results must be taken into consideration to establish diagnosis.

A positive test does not rule out the possibility that other antibiotic resistance mechanisms may be present.

## XII. TECHNICAL PROBLEMS/COMPLAINTS

If you encounter a technical problem or if performances do not correspond with those indicated in this package insert:

- Record the kit batch number
- If possible, keep the sample in the appropriate storage condition during the complaint management
- Contact Coris BioConcept ([client.care@corisbio.com](mailto:client.care@corisbio.com)) or your local distributor

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Last update: JUNE 2018

REF	Catalogue number	Manufacturer
IVD	In vitro diagnostic medical device	Temperature limits
Tests	Contains sufficient for <n> tests	LOT
Instructions	Consult instructions for use	Do not reuse
Keep dry	Keep dry	Use by
DIL SPE	Diluent specimen	CONT NaN <sub>3</sub> Contains Sodium azide