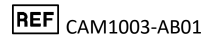


Yersina Pseudotuberculosis Antibody ELISA

An ELISA testkit to detect antibodies against Yersina Pseudotuberculosis in serum or plasma samples





December 2020

Gebruik alleen de juiste versie van het protocol die meegestuurd word met de kit. Please use only the valid version of the package insert provided with the kit. Verwenden Sie nur die jeweils gültige, im Test kit enthaltene, Arbeitsanleitung. Si prega di usare la versione valida dell'inserto del pacco a disposizione con il kit. Por favor, se usa solo la version valida de la metodico técnico incluido aqui en el kit.

Inhoud / Table of Contents / Inhaltsverzeichnis / Tabella die Contenuti / Tabla de Contenidos

| 1. | Introduction | 3 |
|-----|---|----|
| 2. | Intended use of the test kit | |
| 3. | Principle of the test kit | 3 |
| 4. | Contents | 4 |
| 5. | Handling and storage of specimens | 4 |
| 6. | Wash protocol | 5 |
| 7. | Preparations | 5 |
| 8. | Test protocol qualitative | 6 |
| | Before starting this test read "preparations" | 6 |
| 9. | Test protocol quantitative | 8 |
| | Before starting this test read "preparations" | 8 |
| 10. | Precautions | 10 |
| 11. | Validation of the test | 10 |
| 12. | Interpretation of the test results | 11 |
| 13. | Symbols used with EVL ASSAYS | 11 |

1. Introduction

Yersinia pseudotuberculosis is a recognized zoonotic food-borne pathogen. It's a Gram-negative aerobic or facultative anaerobic rod-shaped bacterium in the family *Enterobacteriaceae* and is the causative agent of pseudotuberculosis/rodentiosis, an infectious disease all mammalian and bird species can contract.

For instance, rodents and cats are predisposed. Abscesses can occur in various organs. The pathogen has a high tenacity. In the soil, the pathogen remains infectious for months.

Y. enterocolitica causes enterocolitis. Immunopathological reactions can lead to arthritis, arthrosis and skin diseases. Animals often act as pathogen reservoir.

The infection manifests itself as enteritis, resulting in mucous to bloody diarrhoea.

Intended use of the test kit

The Yersinia Pseudotuberculosis ELISA test kit is designed to detect antibodies against Yersinia Tuberculosis. Purified proteins are attached to the solid phase. After washing the strips are incubated with the sera/plasma to be tested. The strips are washed after incubation to remove unbound materials. A HRPO labeled anti-species conjugate is added to detect bound antibodies to Yersinia Tuberculosis proteins. After incubation and rinsing the substrate is added and the optical density is measured at 450 nm.

3. Principle of the test kit

The principle of the test is based on the reaction of Yersinia Tuberculosis proteins with polyclonal antibodies. To this end, Yersinia Tuberculosis proteins have been coated to a 96 well microtiter plate.

Qualitative

The sample is added (diluted 1:50) to the wells of the coated plate.

Ouantitative

The sample also can be titrated using a 3-step dilution, starting with a dilution 1:50 (\rightarrow 1:150 \rightarrow 1:450 \rightarrow 1:1350).

After washing, the bound avium antibodies are detected by a HRPO conjugated anti-species conjugate.

Color reaction in the wells is directly related to the concentration of Yersina Tuberculosis antibodies in the serum sample.

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4. Contents

- 12 x 8 Microtiter strips coated with purified antigen.
- 1 x Strip holder
- 1 x 18 ml ELISA buffer (green cap)
- 1 x 12 ml HRPO conjugated anti-species antibody (red cap)
- 1 x 0,5 ml Weak Positive control (ready to use) (yellow cap)
- 1 x 1,0 ml Negative control (freeze dried) (silver cap)
- 1 x 20 ml Wash solution (200x concentrated) (black cap), dilute in de-ionized water before use!
- 1 x 8 ml Substrate A (white cap)
- 1 x 8 ml Substrate B (blue cap)
- 1 x 8 ml Stop solution (yellow cap)
- 1 x Plastic cover seal
- 1 x User's manual

Supplies needed (not included)

- Round-bottomed microtiter plate
- Validated precision pipettes
- Pipette tips and clean containers/tubes (EVL)
- ELISA plate reader

5. Handling and storage of specimens

The kit should be stored at 4°C.

An open packet should be used within 10 days.

Samples may be used fresh or may be kept frozen below -20°C before use.

After first use ready-to-use controls and/or reconstituted controls should be aliquoted immediately and stored at -20°C.

Avoid repeated freezing and thawing as this increases non-specific reactivity.

6. Wash protocol

In ELISA's, un-complexed components must be removed efficiently between each incubation step. This is accomplished by appropriate washing. It should be stressed that each washing step must be carried out with care to guarantee reproducible inter- and intra-assay results. It is essential to follow the washing procedures outlined below. Washing may be done manually or with automatic equipment. Automatic washing equipment usually gives better result.

Manual washing

- 1. Empty each well by turning the microtiter plate upside down, followed by a firm vertical downward movement to remove the buffer.
- 2. Fill all the wells with 250 μ l wash solution.
- 3. This washing cycle (step 1 and 2) should be carried out at least 5 times.
- 4. Turn the plate upside down and empty the wells with a firm vertical movement.
- 5. Place the inverted plate on absorbent paper towels and tap the plate firmly to remove any residual wash solution in the wells.
- 6. Take care that none of the wells dry out before the next reagent is added.

Washing with automatic equipment

When automatic plate washing equipment is used, check that all wells are aspirated completely and that the wash solution is correctly added, reaching the rim of each well during each rinsing cycle. The washer should be programmed to execute at least 5 washing cycles.

7. Preparations

- Before using the reagents needed, take them out of the kit and place them on the table for ± 15 min. at room temperature (± 21°C) without exposing them to direct sunlight or (other) heat sources.
- Buffer, controls, standards and conjugates need to be shaken gently before use, in order to dissolve/ mix any components that may have precipitated. Gently tap the vials onto the table, so any fluid still retained in the cap falls back into the solution.
- If fluids need to be mixed into the test well, gently shake by tapping the wells with the fingers or re-suspend with the last pipette tip used for that particular well. Avoid contamination through spattering and prevent any fluid to enter inside the pipette itself.
- Place the reagents back at 4-8°C immediately after use.



8. Test protocol qualitative

Before starting this test read "preparations"

1. Open the packet of strips and take out the strips to be used. Cover the remaining strips with a part of the provided seal and store them at +4°C. and use them within 10 days.

Wash microtiter strip(s) with washing solution, according to washing protocol.

The washing solutions provided must be diluted 200x in aquabidest (5 mega Ohm) water !

Use validated precision pipettes and use a clean pipette tip **before** pipetting the buffer, control, samples, conjugate and substrate.

- 2. Reconstitute directly before use the **negative control** (silver cap) in 1,0ml aquabidest (5 mega Ohm water), divide into aliquots, and store after complete dissolving immediately at -20 °C until use, avoid freeze and thaw cycles.
- 3. Make a three-step dilution of the **positive control** (yellow cap) in **ELISA buffer** (green cap) starting undiluted \rightarrow 1:3 \rightarrow 1:9 \rightarrow 27 in a round-bottomed plate (not supplied).

Example: - Add 180μl positive control to the well 1A.

- Add 120µl ELISA buffer to all other wells 1B, 1C, 1D.
- Transfer 60µl from well 1A to well 1B.
- Mix well and transfer 60µl from well 1B to the well 1C.
- Mix well and transfer 60µl from 1C to the well 1D.
- Mix well and discard 60µl.
- Dilute the negative control (silver cap) 1:50 in ELISA buffer (green cap) in a roundbottomed plate (not supplied).

Example: - Add 147μl ELISA buffer to **well 1E**, add 3μl of the negative control to the **well 1E** and mix well.

Dilute the sample 1:50 in ELISA buffer (green cap) in a round-bottomed plate (not supplied).

Example: - Add 147μl ELISA buffer to **row 1F**, add 3μl of the sample to the **well 1F** and mix well.

- 6. Take 2 wells as substrate controls add only 120µl ELISA buffer (green cap) to these wells.
- 7. Transfer $100\mu l$ of all dilutions to the virus-coated microtiter strips, including the substrate controls.
- 8. Seal and incubate for 60 min at 37°C.
- 9. Wash the strips 5x according to the wash protocol see sub 6.
- 10. Add 100μl HRPO conjugated anti-species antibodies to all wells.
- 11. Seal and incubate for 60 min at 37°C.
- 12. Wash the strips according to the wash protocol see sub 6.
- 13. Mix equal parts of substrate A (white cap) and substrate B (blue cap) with gentle shaking.

 Prepare immediately before use! Only prepare amount needed. Substrate can only be used for 1-2 hours after being mixed.
- 14. Add 100µl substrate solution to each well.
- 15. Incubate 10-15min.in the dark (e.g. cover the wells with a sheet of paper) at room temperature (21°C.). Make sure the negative does not become too dark.

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16. Add **50μl stop solution** to each well; mix well.

Read the absorbency values immediately (within 10 min!) at 450 nm using 620nm as reference on the ELISA reader. <u>Use the substrate controls as blank.</u>



NB: if you pipet directly into the coated ELISA plate with only a small number of samples (<6), make sure the first dilution is done in round bottom microtiter plate second step can be done directly in the coated Elisa plate.

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9. Test protocol quantitative

Before starting this test read "preparations"

1. Open the packet of strips and take out the strips to be used. Cover the remaining strips with a part of the provided seal and store them at +4°C. and use them within 10 days.

Wash microtiter strip(s) with washing solution, according to washing protocol.

The washing solutions provided must be diluted 200x in aquabidest (5 mega Ohm) water !

Use validated precision pipettes and use a clean pipette tip **before** pipetting the buffer, control, samples, conjugate and substrate.

- Reconstitute directly before use the negative control (silver cap) in 1,0ml aquabidest (5 mega Ohm water), divide into aliquots, and store after complete dissolving immediately at -20 °C until use, avoid freeze and thaw cycles.
- 3. Make a pre-dilution of the **negative control** (silver cap) **in ELISA buffer** (green cap) in a round-bottomed plate (not supplied).
 - **Example:** Add 90μl ELISA buffer to **row 1A** and add 10μl of the negative control to the well **1A**.
- 4. Make a pre-dilution of **each sample** in **ELISA buffer** (green cap) in a round-bottomed plate (not supplied).
 - **Example:** Add 90μl ELISA buffer to **row 1B** and add 10μl of the sample to the well **1B**.
- 5. Take 2 wells as substrate controls add only 140µl ELISA buffer (green cap) to these wells.
- 6. Add for dilution of the **positive control** (yellow cap) **100μl ELISA buffer** to **1B, 1C, 1D** of the coated microtiter strip.
- 7. Add for dilution of the **negative control** (brown cap) **120μl ELISA buffer** to **row 1E.** And 100μl to **1F, 1G, 1H** of the coated microtiter strip.
- Add for dilution of the samples 120μl ELISA buffer to the other row 2A and 2E.
 And 100μl to 2B, 2C, 2D and 2F, 2G, 2H (depending on the number of samples) of the coated microtiter strip.
- 9. Make a 3-step dilution of the **positive control** (yellow cap) in the coated microtiter strip, starting Undiluted → 1:3 → 1:9 → 1:27.
 - **Example:** Add 150µl positive control to the well **1A** of the microtiter strip.
 - Mix well and transfer 50 µl to the well 1B
 - Mix well and transfer 50µl to the well 1C
 - Mix well and transfer 50µl to the well 1D
 - Mix well and discard 50µl.
- 10. Make a 3-step dilution of the **negative control** (silver cap) in the coated microtiter strip, starting $1:50 \rightarrow 1:150 \rightarrow 1:450 \rightarrow 1:1350$.
 - **Example:** Add 30µl negative control from step 3 to the well **1E** of the microtiter strip.
 - Mix well and transfer 50µl to the next well 1F

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- Mix well transfer 50µl to the next well 1G
- Mix well and transfer 50µl to the well 1H
- Mix well and discard 50µl.

11. Make 3-step dilution of each sample in the coated microtiter strip, starting 1:50 \rightarrow 1:150 \rightarrow 1:450 \rightarrow 1:1350.



- **Example:** Add 30μl of each sample from step 4 to the well **2A and/or 2E** of the microtiter strip.
 - Mix well and transfer 50 µl to the well 2B and/or 2F
 - Mix well and transfer 50µl to the well 2C and/or 2G
 - Mix well and transfer 50µl to the well 2D and/or 2H
 - Mix well and discard 50µl.
- 12. Add 100µl of the substrate control of step 5 to the last 2 wells of the microtiter strip.
- 13. Seal and incubate for 60 min at 37°C.
- 14. Wash the strips according to the wash protocol see sub 6.
- 15. Add 100µl HRPO conjugated anti-species antibodies to all wells.
- 16. Seal and incubate for 60 min at 37°C.
- 17. Wash the strips according to the wash protocol see sub 6.
- 18. Mix equal parts of buffer A (white cap) and buffer B (blue cap) with gentle shaking.

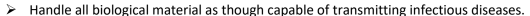
 Prepare immediately before use! Only prepare amount needed. Substrate can only be used for 1-2 hours after being mixed.
- 19. Add 100µl substrate solution to each well.
- 20. Incubate 10-20 min. in the dark (e.g. cover the wells with a sheet of paper) at room temperature (21°C.). Make sure the negative control does not become too dark.
- 21. Add 50µl stop solution to each well; mix well.
- 22. Read the absorbency values immediately (within 10 min!) at 450 Nm by using an ELISA reader. Use the substrate controls as blank.

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10. Precautions



- Do not pipette by mouth.
- > Do not eat, drink, smoke or prepare foods, or apply cosmetics within the designated working area.
- TMB substrate (buffer B) is toxic by inhalation, through contact with skin or when swallowed; observe care when handling substrate.
- ➤ Do not use components past the expiry date and do not mix components from different serial lots.
- > Optimal, results will be obtained by strict adherence to this protocol. Careful pipetting and washing throughout this procedure are necessary to maintain precision and accuracy.
- Each well is ultimately used as an optical cuvette. Therefore, do not touch the under-surface of the microtiter plate and protect it from damage and dirt.

11. Validation of the test

Qualitative:

- > The results are valid if the following criteria are met:
 - The mean value (MV) of the measured OD value for the Positive Control (PC), diluted
 1:3. must be ≥0.800.
 - \circ The MV of the measured OD value for the Negative Control (NC), diluted 1:50, must be ≤0.200.

In case of invalid assays the test should be repeated after a thorough review of the instructions for use.

Calculation

Calculate the mean values (MV) of the measured OD for the Negative Control (NC) and the Positive Control (PC).

The ratio (S/P) of sample OD to mean OD of the positive control is calculated according to the following equation:

$$S/P = \frac{OD_{sample} - MV OD_{NC}}{MV OD_{PC} - MV OD_{NC}}$$

Quantitative:

In order to confirm appropriate test conditions the OD of the positive control, undiluted, should be \geq 0.800 OD units (450nm) and give an endpoint titer of \geq 9.

The negative control, diluted 1:50, should be lower than 0.250 OD units (450nm) and give an endpoint titer of \leq 50.

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Interpretation of the test results 12.

This test can be used in 2 ways.

Qualitative: Positive – Negative

- ➤ A sample with the S/P ratio <0.34 is negative.
 - o Specific antibodies to Yersina Pullorum could not be detected.
- ➤ A sample with the S/P ratio ≥0.34 is positive.
 - o Specific antibodies to Yersina Pullorum were detected.

Quantitative: End point titre

- The ELISA titre can be calculated by constructing a curve and using cut-off line(dilution 1:50-150-450 -1350-4050 -12150 etc. total 8 dilutions of 3 steps) OD on Y-as and Titre on X-as
- Elisa titres can be calculated using as cut-off 2,5 times OD value of negative control at 1:50.

Symbols used with EVL ASSAYS 13.

| Symbol | English | Deutsch | Français | Español | Italiano |
|----------------|--|-----------------------------------|--|---|---------------------------------------|
| Ti | Consult instructions for use | Gebrauchsanweisung beachten | Consulter les instructions d'utilisation | Consulte las instrucciones de uso | Consultare le istruzioni per l'uso |
| (€ | European Conformity | CE-Konfirmitäts- kennzeichnung | Conformité aux normes européennes | Conformidad europea | Conformità europea |
| IVD | In vitro diagnostic device | In-vitro-Diagnostikum | Usage Diagnostic in vitro | Para uso Diagnóstico in vitro | Per uso Diagnostica in vitro |
| RUO | For research use only | Nur für Forschungszwecke | Seulement dans le cadre de recherches | Sólo para uso en investigación | Solo a scopo di ricerca |
| REF | Catalogue number | Katalog-Nr. | Numéro de catalogue | Número de catálogo | Numero di Catalogo |
| LOT | Lot. No. / Batch code | Chargen-Nr. | Numéro de lot | Número de lote | Numero di lotto |
| \sum | Contains sufficient for <n> tests/</n> | Ausreichend für "n" Ansätze | Contenu suffisant pour "n" tests | Contenido suficiente para <n> ensayos</n> | Contenuto sufficiente per "n" saggi |
| 1 | Storage Temperature | Lagerungstemperatur | Température de conservation | Temperatura de conservación | Temperatura di conservazione |
| | Expiration Date | Mindesthaltbarkeits- datum | Date limite d'utilisation | Fecha de caducidad | Data di scadenza |
| ~ | Legal Manufacturer | Hersteller | Fabricant | Fabricante | Fabbricante |
| Distributed by | Distributor | Vertreiber | Distributeur | Distribuidor | Distributore |
| Content | Content | Inhalt | Conditionnement | Contenido | Contenuto |
| Volume/No. | Volume / No. | Volumen/Anzahl | Volume/Quantité | Volumen/Número | Volume/Quantità |

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