

# RIDA®QUICK ADM Monitoring Control Set

**REF** GP3043





#### 1. Intended use

For in vitro diagnostic use. The RIDA<sup>®</sup>QUICK ADM Monitoring Control Set is intended for the use with the corresponding, batch specific kit RIDA<sup>®</sup>QUICK ADM Monitoring (GN3043).

### 2. Summary and explanation of the test

RIDA<sup>®</sup>QUICK ADM Monitoring Control Set is used to check the test reagents and to perform the RIDA<sup>®</sup>QUICK ADM Monitoring (GN3043) test procedure. RIDA<sup>®</sup>QUICK ADM Monitoring Control Set is sold as a separate accessory for RIDA<sup>®</sup>QUICK ADM Monitoring and must be used with the lot-compliant RIDA<sup>®</sup>QUICK ADM Monitoring (GN3043).

#### 3. Test principle

RIDA<sup>®</sup>QUICK ADM Monitoring Control Set is ready for use and can be used like a serum or plasma specimen.

#### 4. Reagents provided

High control 1.2 ml Lot-specific high positive control

Low control 1.2 ml Lot-specific low positive control

The positive controls contain the therapeutic antibody adalimumab. The high positive control has a concentration of approximately 12  $\mu$ g/ml, and the low positive control a concentration of approximately 5  $\mu$ g/ml. The exact specifications can be found in the respective certificate included with RIDA<sup>®</sup>QUICK ADM Monitoring Control Set. A result within this concentration range indicates proper function of the components and correct processing.

#### 5. Storage instructions

RIDA<sup>®</sup>QUICK ADM Monitoring Control Set must be stored at 2-8 °C and may be used up to the expiration date printed on the product. After the expiration date, the quality guarantee is no longer valid. Non-specific stained or additional bands indicate possible expiration of the reagents.

#### 6. Reagents required but not provided

Not included are RIDA<sup>®</sup>QUICK ADM Monitoring (GN3043) Kit and the required accessories.

#### 7. Warnings and precautions for the users

Indicated for *in vitro* diagnostic use only. This test must be carried out only by trained laboratory personnel. The guidelines for working in medical laboratories must be followed. Always adhere strictly to the user instructions for carrying out this test. The reagents contain sodium azide as a preservative. This substance must not be allowed to come into contact with skin or mucous membranes.

## 8. Test procedure

Before using the product, bring the positive controls, the reagents contained in RIDA®QUICK ADM Monitoring (GN3043) Kit, and the test strips to room temperature (20 - 25 °C).

First dilute the high positive control High control and the low positive control Low control in sample diluent Sample diluent 1:50 (20  $\mu$ l specimen + 980  $\mu$ l sample diluent). In a separate reaction vial, mix 90  $\mu$ L Reagent | A (blue liquid, bottle with blue lid) with 90  $\mu$ L Reagent | B (yellow liquid, bottle with transparent lid). Pipette 20  $\mu$ L of the diluted specimen solution into the 180  $\mu$ L of the mixture of reagents A and B, which is equivalent to a further dilution of the sample of 1:10. Incubate at room temperature for 5 minutes, then apply 100  $\mu$ L of each positive control diluted to 1:500 to a separate test cassette.

Incubate for 15 minutes. During that time, a band appears at the level of the test line labeled "T". Another band appears at the level of the control line labeled "C".

## 9. Quality control - indication of instability or deterioration of reagents

The test can only be evaluated, if the test cassette is unharmed and there are no color changes or lines present before applying the sample suspension. The control line (marked with C on the test cassette) has to show up in every test run. In case this band is missing, the following should be checked before repeating the test:

- Expiry date of the reagents and test cassette used
- Correct test procedure
- Contamination of reagents

If the control line is still not visible after repeating the test with a different test cassette contact the manufacturer or your local R-Biopharm distributor.

#### 10. Evaluation and interpretation

The read out is performed on the RIDA<sup>®</sup>QUICK SCAN II (also see RIDA<sup>®</sup>QUICK SCAN II-manual).

The intensity of the test bands depends on the concentration of adalimumab in the control samples used and on the incubation period. Only after the total run time of 15 minutes the final test result can be determined by using the RIDA<sup>®</sup>QUICK SCAN II.

The bands can change during the total incubation time and may also change after drying. The color of the band can vary from red to blue-violet/grey.

## 11. Version history

Version number	Chapter and description
2018-06-28	Release document

## 12. Explanation of symbols

## General symbols

For in vitro diagnostic use

Lot number

Manufacturer

## Test-specific symbols

High control Lot-specific high positive control

Low control Lot-specific low positive control