

BIOLABO www.biolabo.fr **MANUFACTURER: BIOLABO SAS,**

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TECHNICAL SUPPORT AND ORDERS

RF Standard Set

For calibration of quantitative determination of Rheumatoid Factor by Turbidimetric immunoassay

REF RF CALSET51 R1 1x 1 mL, R2 1x 1 mL, R3 1x 1 mL, R4 1x 1 mL, R5 1x 1 mL

 $C \in$

IVD

Made in France

I: corresponds to significant modifications

Latest revision: www.biolabo.fr

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INTENDED USE

Standard Set for the preparation of reference curve for quantitative immunochemical determination of RF in human serum.

Suitable for manual procedure or automated instruments with BIOLABO reagents:

REF RF050E, RF520E, REF K1RF1, K2RF1, K4RF1

R	Ε	Α	G	Ε	N	TS

R1	RF1	⊗	
R2	RF2	(32)	
R3	RF3	Human origin	
R4	RF4		
R5	RF5		

5 vials of RF Standards (5 different levels)

Liquid stabilized plasmas supplemented with RF and diluted in saline

SAFETY CAUTIONS (1) (2)

BIOLABO reagents are designated for professional use in laboratory.

- Material Safety Data Sheet is available upon request.
- Each donor unit used to manufacture this product was tested and found non-reactive for HbsAg, antibody to Hepatitis C and antibody to HIV-1/HIV-2.
- However, no test method can offer complete assurance that infectious agents are absent. All specimens or reagents from biological origin should be handled as potentially infectious, in accordance with good laboratory practices using appropriate precautions.
- Waste disposal: Respect legislation in force in the country.

Any serious incident that has occurred in connection with the device is notified to the manufacturer and the competent authority of the Member State in which the user and/or patient is based.

REAGENTS PREPARATION

Ready for use

I MATERIAL REQUIRED BUT NOT PROVIDED

- 1.BIOLABO Reagents (§ INTENDED USE)
- 2. REF RF CONT1: RF Control

QUALITY CONTROL

Verify the integrity of each vials and batch-specific values before use. Run in accordance with the IFU of the reagent used.

STABILITY AND STORAGE

Stored away from light, well cap in the original vial at 2-8°C. standards are stable when stored and used as described:

• Until the expiry date stated on the label of the Kit.

Once opened:

- Transfer requested quantity, well recap vials and store at 2-8°C.
- Well recapped in the original vial, at least for 6 weeks when free from contamination.

Do not freeze

PROCEDURE

Run in accordance with the IFU of the reagent used.

CALIBRATION VALUES (3)

- Values are traceable to a reference material Standardisation)
- Batch-specific values are indicated on the label of each vial

LIMITS

Factors which may influence results are bacterial contamination, accuracy of reconstitution volume, respect of automated instrument procedure, temperature...

REFERENCES

- Occupational Safety and Health Standards; Bloodborne pathogens (29CFR1910.1030) Federal Register July 1, (1998); 6, p.267-280
- Directive du conseil de l'Europe (90/679/CEE) J. O. de la communauté européenne n°L374 du 31.12.1990, p.1-12
- TIETZ N.W. Textbook of clinical chemistry, 3rd Ed. C.A. Burtis, E.R. Ashwood, W.B. Saunders (1999) p.520

