#### 0020764574322c501V9.0 **RF-II** Rheumatoid Factors II Order information

REF	CONTENT		Analyzer(s) on which cobas c pack(s) can be used
<b>20764574</b> 322	Rheumatoid Factors II 100 tests	System-ID 07 6457 4	cobas c 311, cobas c 501/502
12172828 322	Preciset RF (5 x 1 mL)	Codes 725-729	
<b>03005496</b> 122	RF Control Set (4 x 1 mL)	Code 215 Level I	
		Code 216 Level II	

04489357 190 Diluent NaCl 9 % (50 mL)

#### English

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System information

For **cobas c** 311/501 analyzers: **RF-II:** ACN 017 For **cobas c** 502 analyzer: **RF-II:** ACN 8017

#### Intended use

In vitro test for the quantitative determination of Rheumatoid Factors (RF-II) in human serum and plasma on Roche/Hitachi **cobas c** systems. Measurements may be used as an aid in the diagnosis of rheumatoid arthritis.

#### Summary<sup>1,2,3,4,5,6,7,8,9,10</sup>

Rheumatoid factors are a heterogeneous group of autoantibodies directed against the antigenic determinants on the Fc-region of IgG molecules. They are important in the diagnosis of rheumatoid arthritis, but can also be found in other inflammatory rheumatic diseases and in various non-rheumatic diseases. They are also found in clinically healthy persons over 60 years of age. Despite these restrictions, the detection of rheumatoid factors is a diagnostic criterion of the American College of Rheumatology for classifying rheumatoid arthritis. The autoantibodies occur in all the immunoglobulin classes, although the usual analytical methods are limited to the detection of rheumatoid factors of the IgM type.

The classic procedure for the quantitation of rheumatoid factors is by agglutination with IgG-sensitized sheep erythrocytes or latex particles. Particular problems of these semiquantitative methods are the poor between-laboratory precision and reproducibility, together with standardization difficulties. For these reasons, new assay methods such as nephelometry, turbidimetry, enzyme-immunoassays and radioimmunoassays have been developed. The Roche RF assay is based on the immunological agglutination principle with enhancement of the reaction by latex.

#### Test principle<sup>4,5,6</sup>

Immunoturbidimetric assay.

Latex-bound heat-inactivated IgG (antigen) reacts with the RF-antibodies in the sample to form antigen/antibody complexes which, following agglutination, are measured turbidimetrically.

#### **Reagents - working solutions**

- R1 Glycine buffer: 170 mmol/L, pH 8.0; polyethylene glycol: 0.05 %; bovine serum albumin; stabilizer; preservative
- R2 Latex particles coated with human IgG; glycine buffer: 170 mmol/L, pH 7.3; stabilizer; preservative

#### R1 is in position B and R2 is in position C.

#### Precautions and warnings

#### For in vitro diagnostic use.

Exercise the normal precautions required for handling all laboratory reagents.

Disposal of all waste material should be in accordance with local guidelines. Safety data sheet available for professional user on request.

All human material should be considered potentially infectious. All products derived from human blood are prepared exclusively from the blood of donors tested individually and shown to be free from HBsAg and antibodies to HCV and HIV. The testing methods used assays approved by the FDA or cleared in compliance with the European Directive 98/79/EC, Annex II, List A.

However, as no testing method can rule out the potential risk of infection with absolute certainty, the material should be handled with the same level

System-ID 07 6869 3

of care as a patient specimen. In the event of exposure, the directives of the responsible health authorities should be followed.  $^{\rm 11,12}$ 

For USA: Caution: Federal law restricts this device to sale by or on the order of a physician.

This kit contains components classified as follows in accordance with the Regulation (EC) No. 1272/2008:

2-methyl-2H-isothiazol-3-one hydrochloride

EUH 208 May produce an allergic reaction.

Product safety labeling follows EU GHS guidance.

#### Reagent handling

Ready for use

Mix **cobas c** pack well before placing on the analyzer.

Carefully invert reagent container several times prior to use to ensure that the reagent components are mixed.

#### Storage and stability

RF-II

Shelf life at 2-8 °C:	See expiration date on <b>cobas c</b> pack label.
On-board in use and refrigerated on the analyzer:	8 weeks
Diluent NaCl 9 %	
Shelf life at 2-8 °C:	See expiration date on <b>cobas c</b> pack label.

On-board in use and refrigerated on the analyzer: 12 weeks

#### Specimen collection and preparation

For specimen collection and preparation only use suitable tubes or collection containers.

Only the specimens listed below were tested and found acceptable. Serum

Plasma: Li-heparin and K<sub>2</sub>-EDTA plasma

The sample types listed were tested with a selection of sample collection tubes that were commercially available at the time of testing, i.e. not all available tubes of all manufacturers were tested. Sample collection systems from various manufacturers may contain differing materials which could affect the test results in some cases. When processing samples in primary tubes (sample collection systems), follow the instructions of the tube manufacturer.

Centrifuge samples containing precipitates before performing the assay. See the limitations and interferences section for details about possible sample interferences.

Stability:13	
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8 days at 4-8 °C

1 day at 20-25 °C

3 months at -20 °C (freeze only once)

Sample stability claims were established by experimental data by the manufacturer or based on reference literature and only for the temperatures/time frames as stated in the method sheet. It is the responsibility of the individual laboratory to use all available references and/or its own studies to determine specific stability criteria for its laboratory.



#### Materials provided

See "Reagents - working solutions" section for reagents.

#### Materials required (but not provided)

- See "Order information" section
- General laboratory equipment

#### Assay

For optimum performance of the assay follow the directions given in this document for the analyzer concerned. Refer to the appropriate operator's manual for analyzer-specific assay instructions.

The performance of applications not validated by Roche is not warranted and must be defined by the user.

#### Application for serum and plasma

cobas c 311 test definition			
Assay type	2-Point End		
Reaction time/Assay points	10 / 7-18		
Wavelength (sub/main)	800/570 nm		
Reaction direction	Increase		
Unit	IU/mL		
Reagent pipetting		Diluent (H <sub>2</sub> O)	)
R1	90 µL	-	
R2	30 µL	-	
Sample volumes	Sample	Samp	le dilution
		Sample	Diluent (NaCl)
Normal	3 µL	-	-
Decreased	6 µL	15 µL	135 µL
Increased	3 µL	_	_

#### cobas c 501 test definition

Assay type	2-Point End		
Reaction time/Assay points	10 / 12-26		
Wavelength (sub/main)	800/570 nm		
Reaction direction	Increase		
Unit	IU/mL		
Reagent pipetting		Diluent (H <sub>2</sub> C	))
R1	90 µL	-	
R2	30 µL	-	
Sample volumes	Sample	Sam	ole dilution
		Sample	Diluent (NaCl)
Normal	3 µL	-	-
Decreased	6 µL	15 µL	135 µL

#### cobas c 502 test definition

Increased

Assay type	2-Point End		
Reaction time/Assay points	10 / 12-26		
Wavelength (sub/main)	800/570 nm		
Reaction direction	Increase		
Unit	IU/mL		
Reagent pipetting		Diluen	t (H <sub>2</sub> O)
R1	90 µL	-	
R2	30 µL	-	
Sample volumes	Sample		Sample dilution

3μL

		Sample	Diluent (NaCl)
Normal	3 µL	-	-
Decreased	6 µL	15 µL	135 µL
Increased	6 µL	-	-
Calibration			
Calibrators	S1: H <sub>2</sub> O		
	S2-6: Pree	ciset RF	
Calibration mode	RCM		
Calibration frequency	Full calibra • after 180 • after rea • as required procedure	ation ) days during she gent lot change red following qua s	elf life lity control
Calibration interval may be calibration by the laboratory	extended bas y.	sed on acceptabl	e verification of
Standard 64/2.	nas been sta	indardized using	the WHO
<b>Quality control</b> For quality control, use con section.	trol materials	as listed in the "	Order information"
In addition, other suitable c	ontrol materia	al can be used.	
The control intervals and lir individual requirements. Va limits. Each laboratory shou values fall outside the defin	mits should be dues obtained uld establish o ned limits.	e adapted to each I should fall within corrective measu	n laboratory's n the defined res to be taken if
-ollow the applicable gover quality control.	rnment regula	tions and local g	uidelines for
Calculation			
cobas c systems automatic sample.	cally calculate	e the analyte con	centration of each
Limitations - interference			
Criterion: Recovery within ± 14 IU/mL.	± 10 % of initi	al value at an RF	concentration of
Icterus: <sup>15</sup> No significant inte and 60 for unconjugated bil concentration: 624 µmol/L bilirubin concentration: 102	erference up t lirubin (appro: or 40 mg/dL) 6 µmol/L or 6	o an I index of 40 ximate conjugate and approximate 0 mg/dL).	) for conjugated d bilirubin unconjugated
Hemolysis: <sup>15</sup> No significant	interference	up to an H index	of 300 00 mg/dL)
Lipemia (Intralipid): <sup>15</sup> No signature is poor correlation be	gnificant inter etween the L i	ference up to an index (correspond	L index of 2000. ds to turbidity) and
Drugs: No interference was	s found at the	rapeutic concentr	rations using
High dose hook-effect: Usir flag was observed up to an	ng the prozon	e check, no false ation of 6000 IU/r	e result without a mL.
n very rare cases, gammo nacroglobulinemia), may c	pathy, in parti ause unreliab	cular type IgM (V le results. <sup>18</sup>	Valdenström's
There is the possibility that with the test and cause unr	other substar eliable results	nces and/or facto 3.	rs may interfere
For diagnostic purposes, th conjunction with the patient findings.	t's medical his	uld always be as story, clinical exa	sessed in mination and other
ACTION REQUIRED Special Wash Programmi when certain test combinat latest version of the carry-o SMS-SmpCln1+2-SCCS M operator's manual. cobas o necessary for avoiding carr input is required in certain o	ing: The use ions are run t over evasion li lethod Sheets c 502 analyze y-over is avai cases.	of special wash s ogether on <b>coba</b> ist can be found v . For further instr or: All special was lable via the <b>cob</b>	steps is mandatory <b>s c</b> systems. The with the NaOHD- ructions refer to the sh programming <b>has</b> link, manual

Where required, special wash/carry-over evasion programming must be implemented prior to reporting results with this test.

#### 2/4

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### 0020764574322c501V9 0 Rheumatoid Factors II

#### Limits and ranges

#### Measuring range

#### 10-130 IU/mL

Determine samples having higher concentrations via the rerun function. Dilution of samples via the rerun function is a 1:5 dilution. Results from samples diluted using the rerun function are automatically multiplied by a factor of 5.

#### Lower limits of measurement

Lower detection limit of the test

10 IU/mL

The lower detection limit represents the lowest measurable analyte level that can be distinguished from zero. It is calculated as the value lying 3 standard deviations above that of the lowest standard (standard 1 + 3 SD, repeatability, n = 21).

#### Expected values

< 14 IU/mL

This value is based on serum samples from 541 test subjects.

Each laboratory should investigate the transferability of the expected values to its own patient population and if necessary determine its own reference ranges

#### Specific performance data

Representative performance data on the analyzers are given below. Results obtained in individual laboratories may differ.

#### Precision

Precision was determined using human samples and controls in an internal protocol with repeatability (n =  $\tilde{2}1$ ) and intermediate precision (3 aliquots per run, 1 run per day, 21 days). The following results were obtained:

Mean	SD	CV
IU/mL	IU/mL	%
23.7	0.2	0.8
53.0	0.5	0.9
19.5	0.3	1.6
27.5	0.3	1.1
Mean	SD	CV
IU/mL	IU/mL	%
23.2	0.3	1.4
51.4	0.8	1.5
19.3	0.3	1.6
26 1	0.5	10
	Mean IU/mL 23.7 53.0 19.5 27.5 Mean IU/mL 23.2 51.4 19.3 26.1	Mean SD   IU/mL IU/mL   23.7 0.2   53.0 0.5   19.5 0.3   27.5 0.3   Mean SD   IU/mL IU/mL   23.2 0.3   51.4 0.8   19.3 0.3

#### Method comparison

RF values for human serum and plasma samples obtained on a cobas c 501 analyzer (y) were compared with those determined using the corresponding reagent on a Roche/Hitachi 917 analyzer (x).

Sample size (n) = 70

Passing/Bablok <sup>19</sup>	Linear regression
y = 1.000x - 1.20 IU/mL	y = 0.999x - 1.39 IU/mL
т = 0.959	r = 0.998

The sample concentrations were between 10.8 and 114 IU/mL.

#### References

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A point (period/stop) is always used in this Method Sheet as the decimal separator to mark the border between the integral and the fractional parts of a decimal numeral. Separators for thousands are not used.

#### Symbols

Roche Diagnostics uses the following symbols and signs in addition to those listed in the ISO 15223-1 standard (for USA: see dialog.roche.com for definition of symbols used):

CONTENT
GTIN

Contents of kit Volume after reconstitution or mixing Global Trade Item Number

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