



Rheumatoid Factor (RF)

Current Revision and Date ^a	Rev. 02, 2019-11	
Product Name	Atellica CH Rheumatoid Factor (RF)	REF 11097618 (360 tests)
Abbreviated Product Name	Atellica CH RF	
Test Name/ID	RF	
Systems	Atellica CH Analyzer	
Materials Required but Not Provided	Atellica CH LSP CAL	REF 11099434
Specimen Types	Serum	
Sample Volume	11.5 µL	
Measuring Interval	3.5–90.0 IU/mL	

^a A vertical bar in the page margin indicates technical content that differs from the previous version.



Intended Use

The Atellica® CH Rheumatoid Factor (RF) assay is for *in vitro* diagnostic use in the quantitative determination of rheumatoid factor in human serum using the Atellica® CH Analyzer. Such measurements are used in the diagnosis of rheumatoid arthritis. Rheumatoid factors are autoantibodies directed against the Fc portion of IgG. The majority of rheumatoid factors are IgM, but may be IgG or IgA. Conditions giving rise to such factors include rheumatic conditions and chronic inflammatory processes.^{1,2}

Summary and Explanation

The Atellica CH Rheumatoid Factor (RF) assay measures rheumatoid factors in serum by a latex-enhanced immunoturbidimetric assay. It is based on the principle that the analyte concentration is a function of the intensity of scattered light caused by the latex aggregates. The latex particles coated with the antibody rapidly agglutinate in the presence of the rheumatoid factors, forming aggregates.

Principles of the Procedure

The Atellica CH RF Reagent 2 is a suspension of uniform polystyrene latex particles coated with human IgG. When serum containing rheumatoid factor is mixed with the Reagent 2, agglutination takes place resulting in an increase in the turbidity. This turbidity can be measured at 571/805 nm. The rheumatoid factor concentration in serum is determined from a calibration curve that is generated with a set of calibrators.

Reagents

Material Description	Storage	Stabilitya
Atellica CH RF	Unopened at 2–8°C	Until expiration date on product
Pack 1 (P1) Well 1 (W1) Reagent 1 (R1) 9.3 mL 3-N-Morpholino propane sulfonic acid buffer (pH 7.4) (25 mmol/L); sodium azide (0.09%); bovine serum albumin (1.0%)	Onboard per well	21 days
Well 2 (W2) Reagent 1 (R1) 9.3 mL 3-N-Morpholino propane sulfonic acid buffer (pH 7.4) (25 mmol/L); sodium azide (0.09%); bovine serum albumin (1.0%)		
Pack 2 (P2)		
Well 1 (W1) Reagent 2 (R2) 5.0 mL IgG synthetic latex (human); sodium azide (0.09%)		
Well 2 (W2) Reagent 2 (R2) 5.0 mL IgG synthetic latex (human); sodium azide (0.09%)		

^a Refer to Storage and Stability.

Warnings and Precautions

For in vitro diagnostic use.

For Professional Use.

CAUTION

Federal (USA) law restricts this device to sale by or on the order of a licensed healthcare professional.

Safety data sheets (SDS) available on siemens.com/healthineers.



CAUTION POTENTIAL BIOHAZARD

Contains human source material. Each donation of human blood or blood component was tested by FDA-approved methods for the presence of antibodies to human immunodeficiency virus type 1 (HIV-1) and type 2 (HIV-2), as well as for hepatitis B surface antigen (HBsAg) and antibody to hepatitis C virus (HCV). The test results were negative (not repeatedly reactive). No test offers complete assurance that these or other infectious agents are absent; this material should be handled using good laboratory practices and universal precautions.³⁻⁵

CAUTION

This device contains material of animal origin and should be handled as a potential carrier and transmitter of disease.

Contains sodium azide as a preservative. Sodium azide can react with copper or lead plumbing to form explosive metal azides. On disposal, flush reagents with a large volume of water to prevent buildup of azides. Disposal into drain systems must be in compliance with prevailing regulatory requirements.

Dispose of hazardous or biologically contaminated materials according to the practices of your institution. Discard all materials in a safe and acceptable manner and in compliance with prevailing regulatory requirements.

Note For information about reagent preparation, refer to *Preparing the Reagents* in the *Procedure* section.

Storage and Stability

Unopened reagents are stable until the expiration date on the product when stored at 2-8°C. Do not use products beyond the expiration date printed on the product labeling.

Onboard Stability

Reagents are stable onboard the system for 21 days per well. Discard reagents at the end of the onboard stability interval. Do not use products beyond the expiration date printed on the product labeling.

Specimen Collection and Handling

Human serum is the recommended sample type for this assay.

Collecting the Specimen

- Observe universal precautions when collecting specimens. Handle all specimens as if they are capable of transmitting disease.⁵
- Follow recommended procedures for collection of diagnostic blood specimens by venipuncture.⁶
- Follow the instructions provided with your specimen collection device for use and processing.⁷
- Allow blood specimens to clot completely before centrifugation.⁴
- Keep tubes capped at all times.⁴

Storing the Specimen

Separated specimens may be stored for up to 7 days at 2–8°C. Specimens may be stored frozen for up to 3 months at -20°C.⁴

The handling and storage information provided here is based on data or references maintained by the manufacturer. It is the responsibility of the individual laboratory to use all available references and/or its own studies when establishing alternate stability criteria to meet specific needs.

Transporting the Specimen

Package and label specimens for shipment in compliance with applicable federal and international regulations covering the transport of clinical specimens and etiological agents.

Preparing the Samples

This assay requires 11.5 μ L of sample for a single determination. This volume does not include the unusable volume in the sample container or the additional volume required when performing duplicates or other tests on the same sample. For information about determining the minimum required volume, refer to the online help.

Note Do not use specimens with apparent contamination.

Before placing samples on the system, ensure that samples are free of:

- Bubbles or foam.
- Fibrin or other particulate matter.

Note Remove particulates by centrifugation according to CLSI guidance and the collection device manufacturer's recommendations.⁴

Note For a complete list of appropriate sample containers, refer to the online help.

Procedure

Materials Provided

The following materials are provided:

REF	Contents	Number of Tests
11097618	Pack 1 (P1) Well 1 (W1) 9.3 mL of Atellica CH RF Reagent 1 Well 2 (W2) 9.3 mL of Atellica CH RF Reagent 1 Pack 2 (P2) Well 1 (W1) 5.0 mL of Atellica CH RF Reagent 2	2 x 180
	Well 2 (W2) 5.0 mL of Atellica CH RF Reagent 2	

Materials Required but Not Provided

The following materials are required to perform this assay, but are not provided:

REF	Description	
	Atellica CH Analyzer ^a	
11099434	Atellica CH LSP CAL (calibrator)	1 x 1.0 mL calibrator level 1 CAL 1 1 x 1.0 mL calibrator level 2 CAL 2 1 x 1.0 mL calibrator level 3 CAL 3 1 x 1.0 mL calibrator level 4 CAL 4 1 x 1.0 mL calibrator level 5 CAL 5 1 x 1.0 mL calibrator level 6 CAL 6 Calibrator lot-specific value sheet CAL LOT VAL
	Commercially available quality control r	naterials

Additional system fluids are required to operate the system: Atellica CH Diluent, Atellica CH Wash, Atellica CH Conditioner, Atellica CH Cleaner, Atellica CH Reagent Probe Cleaner 1, Atellica CH Reagent Probe Cleaner 2, Atellica CH Reagent Probe Cleaner 4, Atellica CH Lamp Coolant, and Atellica CH Water Bath Additive. For system fluid instructions for use, refer to the Document Library.

Assay Procedure

The system automatically performs the following steps:

- 1. For serum, dispenses 50 μ L of primary sample and 200 μ L of Atellica CH Diluent into a dilution cuvette.
- 2. Dispenses 69 μ L of Reagent 1 into a reaction cuvette.
- 3. Dispenses 11.5 µL of pre-diluted sample into a reaction cuvette.

- 4. Measures the absorbance after sample addition.
- 5. Dispenses 23 μ L of Reagent 2 into a reaction cuvette.
- 6. Mixes and incubates the mixture at 37°C.
- 7. Measures the absorbance after Reagent 2 addition.
- 8. Reports results.

Test Duration: 10 minutes

Preparing the Reagents

All reagents are liquid and ready to use.

Preparing the System

Ensure that the system has sufficient reagent packs loaded in the reagent compartment. For information about loading reagent packs, refer to the online help.

Performing Calibration

For calibration of the Atellica CH RF assay, use Atellica CH LSP CAL. Use the calibrators in accordance with the calibrator instructions for use.

Calibration Frequency

I

Perform a calibration if one or more of the following conditions exist:

- When changing lot numbers of primary reagent packs.
- At the end of the lot calibration interval, for a specified lot of calibrated reagent on the system.
- At the end of the pack calibration interval, for calibrated reagent packs on the system.
- When indicated by quality control results.
- After major maintenance or service, if indicated by quality control results.

At the end of the onboard stability interval, replace the reagent pack on the system with a new reagent pack. Recalibration is not required, unless the lot calibration interval is exceeded.

Stability Interval	Days
Lot Calibration	60
Pack Calibration	14
Reagent Onboard Stability	21

For information about lot calibration and pack calibration intervals, refer to the online help.

Follow government regulations or accreditation requirements for calibration frequency. Individual laboratory quality control programs and procedures may require more frequent calibration.

Performing Quality Control

For quality control of the Atellica CH RF assay, use at least two levels (low and high) of the appropriate quality control material of known analyte concentration. Use the quality control material in accordance with the quality control instructions for use.

For the assigned values, refer to the lot-specific value sheet provided. A satisfactory level of performance is achieved when the analyte values obtained are within the expected control range for the system or within your range, as determined by an appropriate internal laboratory quality control scheme. Follow your laboratory's quality control procedures if the results obtained do not fall within the acceptable limits. For information about entering quality control definitions, refer to the online help.

Follow government regulations or accreditation requirements for quality control frequency. Individual laboratory quality control programs and procedures may require more frequent quality control testing.

Taking Corrective Action

If the quality control results do not fall within the assigned values, do not report results. Perform corrective actions in accordance with established laboratory protocol. For suggested protocol, refer to the online help.

Results

Calculation of Results

The system determines the result using the calculation scheme described in the online help. The system reports results in IU/mL (common units).

For information about results outside the specified measuring interval, refer to *Measuring Interval*.

Interpretation of Results

Results of this assay should always be interpreted in conjunction with the patient's medical history, clinical presentation, and other findings.

Limitations

The Atellica CH RF assay is limited to the detection of rheumatoid factor in human serum.

Do not use hemolyzed samples.

The following substances may interfere with the Atellica CH RF assay when present in serum at the concentrations indicated in the table below. The observed bias due to these substances is shown in the table below.

Substance	Substance Test Concentration mg/dL (mmol/L)	Analyte Concentration IU/mL	Percent Bias
Hemoglobin	200 (0.125)	19.3	-13

Expected Values

Reference Interval

A reference interval for healthy adults was established in accordance with CLSI Document EP28-A3c and verified on the Atellica CH Analyzer.⁸

The reference interval for rheumatoid factor is < 14 IU/mL for adults. These data were established on the ADVIA® Chemistry system.⁹

As with all *in vitro* diagnostic assays, each laboratory should determine its own reference interval for the diagnostic evaluation of patient results. Consider these values as guidance only.⁸

Performance Characteristics

Measuring Interval

The Atellica CH RF assay provides results from 3.5–90.0 IU/mL. The system flags all values that are outside the specified measuring interval.

Extended Measuring Interval

An automatic repeat condition for this assay extends the measuring interval to 900 IU/mL for serum. You may configure the system to trigger an automatic repeat. Automatic repeat results will be flagged **Autorepeat**.

Detection Capability

Detection capability was determined in accordance with CLSI Document EP17-A2. ¹⁰ The assay is designed to have a limit of blank (LoB) < limit of detection (LoD) and LoD \le 3.5 IU/mL.

The LoD corresponds to the lowest concentration of rheumatoid factor that can be detected with a probability of 95%. The LoD for the Atellica CH RF assay is 3.2 IU/mL, and was determined using 120 determinations, with 60 blank and 60 low level replicates, and a LoB of 1.6 IU/mL.

Assay results obtained at individual laboratories may vary from the data presented.

Precision

Precision was determined in accordance with CLSI Document EP05-A3.¹¹ Samples were assayed on an Atellica CH Analyzer in duplicate in 2 runs per day for 20 days ($N \ge 80$ for each sample). The following results were obtained:

			Repeata	ability	Designed to be ≤	Within-Lab	Precision	Designed to be ≤
Sample Type	N	Mean IU/mL		CV ^b (%)	CV (%)	SD IU/mL	CV (%)	CV (%)
Serum QC	80	20.8	0.52	2.5	3.5	0.68	3.3	7.0
Serum Pool	80	50.9	0.27	0.5	3.5	0.52	1.0	7.0
Serum Pool	80	71.1	0.40	0.6	3.5	0.69	1.0	7.0

^a Standard deviation.

Assay results obtained at individual laboratories may vary from the data presented.

Assay Comparison

The Atellica CH RF assay is designed to have a correlation coefficient of > 0.950 and a slope of 1.0 ± 0.10 compared to ADVIA Chemistry 1800 RF. Assay comparison was determined using the Deming linear regression model in accordance with CLSI Document EP09-A3.¹² The following results were obtained:

Specimen	Comparative Assay (x)	Regression Equation	Sample Interval	N ^a	r ^b
Serum	ADVIA Chemistry 1800 RF	y = 0.99x + 0.9 IU/mL	4.8-85.6 IU/mL	100	0.993

^a Number of samples tested.

The agreement of the assay may vary depending on the study design, comparative assay, and sample population. Assay results obtained at individual laboratories may vary from the data presented.

b Coefficient of variation.

b Correlation coefficient.

Interferences

Hemolysis, Icterus, and Lipemia (HIL)

The Atellica CH RF assay is designed to have \leq 10% interference from hemoglobin, bilirubin, and lipemia. Interfering substances at the levels indicated in the table below were tested in accordance with CLSI Document EP07-A2 using the Atellica CH RF assay.¹³

Bias is the difference in the results between the control sample (does not contain the interferent) and the test sample (contains the interferent) expressed in percent. Bias > 10% is considered interference. Analyte results should not be corrected based on this bias.

Substance	Substance Test Concentration Common Units (SI Units)	Analyte Concentration IU/mL	Percent Bias
Hemoglobin	150 mg/dL (0.094 mmol/L)	19.3	-9
	1000 mg/dL (0.624 mmol/L)	56.9	2
Bilirubin, conjugated	25 mg/dL (428 μmol/L)	21.1	4
	25 mg/dL (428 μmol/L)	66.7	-1
Bilirubin, unconjugated	25 mg/dL (428 μmol/L)	18.4	3
	25 mg/dL (428 μmol/L)	62.2	0
Lipemia (Intralipid®)	1000 mg/dL (11.3 mmol/L)	18.6	-2
	1000 mg/dL (11.3 mmol/L)	61.1	-1

Assay results obtained at individual laboratories may vary from the data presented.

High-Dose Hook Effect

High RF levels can cause a paradoxical decrease in signal as a result of the high-dose hook effect. In the Atellica CH RF assay, RF levels as high as 450 IU/mL will read > 90 IU/mL.

Standardization

The assay shall be traceable to an internal standard manufactured using highly purified materials.

Assigned values for calibrators are traceable to this standardization.9

Technical Assistance

For customer support, contact your local technical support provider or distributor. siemens.com/healthineers

References

- 1. Anderson SG, Bentzon MW, Houba V, Krag P. International reference preparation of rheumatoid arthritis serum. *Bull World Health Organ*. 1970;42(2):311–318.
- 2. Galvin JP, Looney CE, Leflar CC, et al. Particle enhanced photometric immunoassay systems. In: Nakamura RM, Ditto WR, Tucker ES, eds. *Clinical laboratory assays: New technology and future directions*. New York, NY: Masson Publishing; 1983:73-95.
- 3. Centers for Disease Control. Perspectives in disease prevention and health promotion update: Universal precautions for prevention of transmission of human immunodeficiency virus, hepatitis B virus and other bloodborne pathogens in healthcare settings. *MMWR*. 1988;37(24):377–382, 387–388.
- 4. Clinical and Laboratory Standards Institute. *Procedures for the Handling and Processing of Blood Specimens for Common Laboratory Tests; Approved Guideline—Fourth Edition*. Wayne, PA: Clinical and Laboratory Standards Institute; 2010. CLSI Document GP44-A4.

5. Clinical and Laboratory Standards Institute. *Protection of Laboratory Workers from Occupationally Acquired Infections; Approved Guideline—Fourth Edition*. Wayne, PA: Clinical and Laboratory Standards Institute; 2014. CLSI Document M29-A4.

- 6. Clinical and Laboratory Standards Institute. *Procedures for the Collection of Diagnostic Blood Specimens by Venipuncture; Approved Standard—Sixth Edition*. Wayne, PA: Clinical and Laboratory Standards Institute; 2007. CLSI Document GP41-A6.
- 7. Clinical and Laboratory Standards Institute. *Tubes and Additives for Venous and Capillary Blood Specimen Collection; Approved Standard—Sixth Edition*. Wayne, PA: Clinical and Laboratory Standards Institute; 2010. CLSI Document GP39-A6.
- 8. Clinical and Laboratory Standards Institute. *Defining, Establishing, and Verifying Reference Intervals in the Clinical Laboratory; Approved Guideline—Third Edition*. Wayne, PA: Clinical and Laboratory Standards Institute; 2010. CLSI Document EP28-A3c.
- 9. Data on file at Siemens Healthcare Diagnostics.
- 10. Clinical and Laboratory Standards Institute. *Evaluation of Detection Capability for Clinical Laboratory Measurement Procedures; Approved Guideline—Second Edition*. Wayne, PA: Clinical and Laboratory Standards Institute; 2012. CLSI Document EP17-A2.
- 11. Clinical and Laboratory Standards Institute. *Evaluation of Precision of Quantitative Measurement Procedures; Approved Guideline—Third Edition*. Wayne, PA: Clinical and Laboratory Standards Institute; 2014. CLSI Document EP05-A3.
- 12. Clinical and Laboratory Standards Institute. *Measurement Procedure Comparison and Bias Estimation Using Patient Samples; Approved Guideline—Third Edition*. Wayne, PA: Clinical and Laboratory Standards Institute; 2013. CLSI Document EP09-A3.
- 13. Clinical and Laboratory Standards Institute. *Interference Testing in Clinical Chemistry; Approved Guideline—Second Edition*. Wayne, PA: Clinical and Laboratory Standards Institute; 2005. CLSI Document EP07-A2.

Definition of Symbols

The following symbols may appear on the product labeling:

Symbol	Symbol Title and Description
ŢŢį.	Consult instructions for use
i Rev. 01	Version of instructions for use
i siemens.com/healthcare i siemens.com/document-library	Internet URL address to access the electronic instructions for use
Rev. REVISION	Revision
\triangle	Caution Consult instructions for use or accompanying documents for cautionary information such as warnings and precautions that cannot, for a variety of reasons, be presented on the medical device.
&	Biological risks Potential biological risks are associated with the medical device.
	Corrosive

Symbol	Symbol Title and Description
(Dangerous to environment
(•)	Irritant Oral, dermal, or inhalation hazard
	Inhalation hazard Respiratory or internal health
	Flammable Flammable to extremely flammable
	Oxidizing
	Explosive
	Toxic
\Diamond	Compressed gas
*	Keep away from sunlight Prevent exposure to sunlight and heat.
<u>tt</u>	Up Store in an upright position.
	Do not freeze
1 2°C √ 8°C	Temperature limit Upper and lower limits of temperature indicators are adjacent to the upper and lower horizontal lines.
	Handheld barcode scanner
IVD	In vitro diagnostic medical device
\sum_{Σ} (n)	Contains sufficient for <n> tests Total number of IVD tests the system can perform with the IVD kit reagents appears adjacent to the symbol.</n>
RxOnly	Prescription device (US only) Applies only to United States-registered IVD assays. CAUTION: Federal (USA) law restricts this device to sale by or on the order of a licensed healthcare professional.

Symbol	Symbol Title and Description
2	Mixing of substances Mix product before use.
g AmL	Reconstitute and mix lyophilized product before use.
mL → I ←	Target
← →	Interval
	Legal Manufacturer
EC REP	Authorized Representative in the European Community
Ξ	Use-by date Use by the designated date.
LOT	Batch code
REF	Catalog number
43	Recycle
PRINTED WITH SOY INK	Printed with soy ink
CE	CE Mark
€	CE Mark with notified body ID number Notified body ID number can vary.
YYYY-MM-DD	Date format (year-month-day)
CHECKSUM	Variable hexadecimal number that ensures the Master Curve and Calibrator definition values entered are valid.
UNITS C	Common Units
UNITS SI	International System of Units
MATERIAL	Material
MATERIAL ID	Unique material identification number
CONTROL NAME	Name of control
CONTROL TYPE	Type of control

Legal Information

Atellica and ADVIA are trademarks of Siemens Healthcare Diagnostics.

All other trademarks are the property of their respective owners.

© 2017-2019 Siemens Healthcare Diagnostics. All rights reserved.

Siemens Healthcare Diagnostics Inc. 511 Benedict Avenue Tarrytown, NY 10591 USA siemens.com/healthineers

Siemens Healthineers Headquarters

Siemens Healthcare GmbH Henkestr. 127 91052 Erlangen Germany Phone: +49 9131 84-0

siemens.com/healthineers