

CRP

C-reactive Protein Test Kit
(Nephelometry Immunoassay Method)

Instructions for Use

Version: B/6

REF HP-CRP-25

Manufacturer

 Shijiazhuang HiPro Biotechnology Co.,Ltd.
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Product Name

General Name: C-reactive Protein Test Kit (Nephelometry
Immunoassay Method)

Specification

Package Specification
25 Tests/ Kit.

Intended Use

This product is used to determine the content of C-reactive
proteinTest Kit(CRP) in human blood.

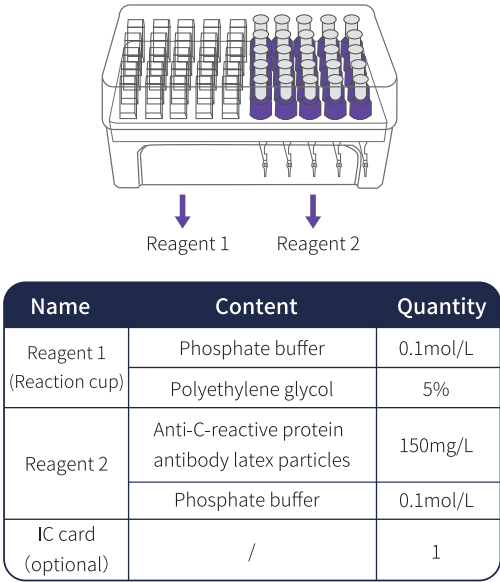
Test Principle

The antibody of C-reactive protein Test Kit is coated on the
latex surface. The CRP in the sample and the antibody become
to immune complexes by Latex agglutination reaction. The
immune complexes will produce the phenomenon of light
scattering which is proportional to the intensity of scattered
light and samples of CRP levels. Using specific protein analyzer
to measure the intensity of scattered light, the concentration
of CRP is determined by comparing the turbidity of samples to
the standard concentration.

Component

The CRP test kit consists of two reagents R1 and R2, as shown
on Figure 1.

Figure 1



Do not mix different batches of reagents.

Storage&stability

Store the test kit at 2°C-8°C until the expiration date indicated
on the label. The test kit is stable for one year when unopened.

Use up the test kit within one month after opening the pack-
age.

Do not freeze the test kit.

Do not mix different lots of the test kit.

Special Instrument Requirements

HP-083/4-I POCT Immunoassay System,
HP-083/4-II POCT Immunoassay System,
HP-AFS/1 Automatic Immunoassay System,
HP-AFS/3 Automatic Immunoassay System.

Specimen type

Whole blood. The sample store at 2-8°C for 48 hours, -20°C for
1 month. Avoid repeated freezing. Before test, ensure fully
mixed.

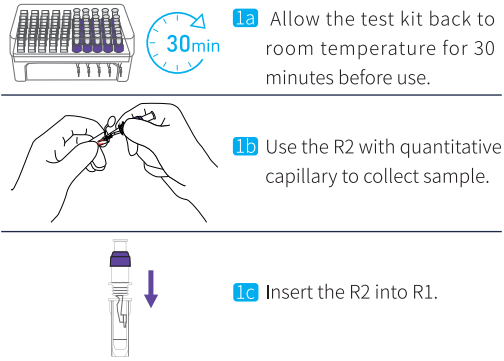
Procedures

HP-083/4-I&HP-083/4-II POCT Immunoassay System

Note:

- Please read user manual of HP-083/4-I and HP-083/4-II
before use;
- The analyzer will finish self check after start-up;
- Insert the IC card of CRP test kit to let analyzer read the
parameter;
- The analyzer calibration can be done with app. It is recom-
mended that analyzer calibration should be done for each
new lot of test kit.

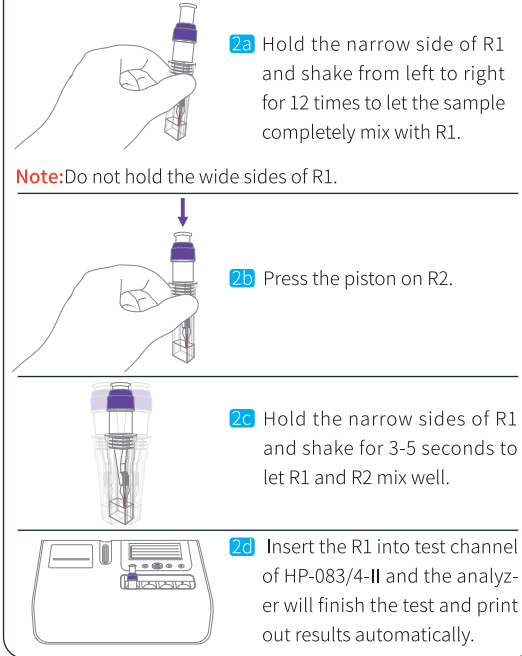
Step 1 Sample Preparation



Note:

- The parameter is built in the IC card.
- Please insert the corresponding IC card into analyzer
to let the analyzer read the parameter before each assay test.
- The capillary of the R2 should be fully filled.

Step 2 Testing

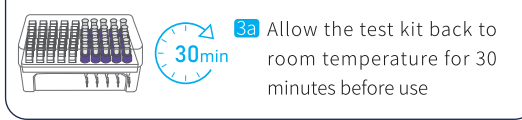


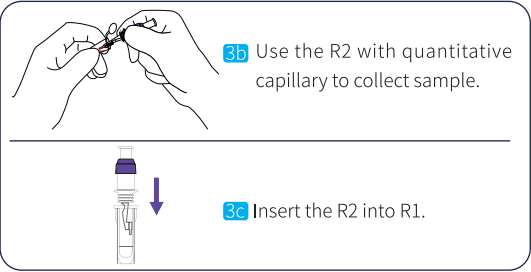
HP-AFS/1&HP-AFS/3 Automatic Immunoassay System

Note:

- Please read user manual of HP-AFS/1 and HP-AFS/3 before
use;
- The analyzer will finish the self check after start-up;
- It is recommend to do analyzer calibration monthly and for
each new lot of test kit.

Step 1 Sample Preparation





- Note:**
- Please update the standard curve with the barcode on the R1 cuvette if a new lot test kit is to be used.
 - The capillary of the R2 should be fully filled.

Step 2 Testing

- 4a** Insert the R1 into test channel of analyzer.
- 4b** The analyzer will mix the sample from R2 capillary with R1 automatically.
- 4c** The analyzer will mix the R2 and R1 automatically.
- 4d** The analyzer will test and print the results automatically.

Calibration

The calibration values for the different lots of the kits are stored on the calibration IC card or the two-dimensional code on the cuvette. Before test the new lot of kits, read the calibration card parameters first. Or the instrument automatically scan the two-dimensional code on the cup to obtain the corresponding calibration curve during testing.

Quality control

3- level calibration system guarantee the results' reliability for each lot of test kits, including the instrument calibration, remote reagent calibration and the third party calibration.

The third party calibration applicable for:

1. The daily indoor quality control test.
2. New lots of reagent.
3. New operator training.
4. The results can not match the clinical symptoms.
5. The first use of the reagent.

If still can not be calibrated, contact the manufacture for further technical support.

Reference Value

Diagnosis of Infection and inflammation for adults, the reference value: CRP<10mg/L

Without inflammation, the risk of cardiovascular disease: CRP <1.0 mg/L.

For newborn, the reference value:

Newborn (Age 0-4 weeks): 0.1-4.1 mg/L

It is important testing the concentration of CRP in the acute phase of the disease, recommended that each laboratory establish its own reference range.

Interpretation

The test results of this reagent are only for clinical reference. the clinical diagnosis and treatment of patients should be considered in combination with their symptoms / signs, medical history, other laboratory tests and treatment responses.

Limitations

1. When bilirubin≥684μmol/L, triglyceride≥15mmol/L it has interference to this determination.
2. This product is based on latex agglutination system, only suitable for the specific analyzer.
3. The sample for myeloma patient may precipitate in R1 to affect the result.
4. The sample for extended periods may lead to differences in test results.

Performance Characteristics

1. Linearity range:0.5 mg/L ~ 200mg/L, r≥0.990
 2. Detection limit: ≤0.3 mg/L
- The limit of detection means the lowest detectable analyte level that can distinguish the concentration. Calculate based on the minimum standard above the two standard deviation of the data (Blank table, 1+2SD, within-run precision, n=20).
3. Precision
- Test the control material by CRP Test Kit 2 times per day for 20 days (n=80) according to EP5-A2 of CLSI.

The data as below:

a.

HP-083/4-II POCT Immunoassay System					
Sample	Mean mg/L	Within-Run		Between-Run	
		S.D.	%C.V.	S.D.	%C.V.
Control 1	21.5	0.96	4.5	1.21	5.6
Control 2	145.6	5.21	3.6	5.36	3.7

b.

HP-AFS/3 Automatic Immunoassay System					
Sample	Mean mg/L	Within-Run		Between-Run	
		S.D.	%C.V.	S.D.	%C.V.
Control 1	22.3	1.15	5.2	1.32	5.9
Control 2	146.3	5.05	3.5	5.63	3.8

c.

HP-AFS/1 Automatic Immunoassay System					
Sample	Mean mg/L	Within-Run		Between-Run	
		S.D.	%C.V.	S.D.	%C.V.
Control 1	22.1	1.32	6.0	1.25	5.7
Control 2	145.2	5.63	3.9	6.32	4.4

4.Methodology comparison

Compared to Hitachi 7060 CRP by test the same sample, the relative data as below:

HP-AFS/3 Automatic Immunoassay System				
Site No.	Sample Type	No.of Assays	Regression Line	Coefficient correlation
1	Whole blood	50	Y= 1.02X+0.64	0.98

The concentration of sample is about 0.6mg/L-197mg/L

Precautions

- ⚠ Attention:**
- Only for in vitro diagnostic.
- Only for professional use.
- All samples and reactive wastes are treated as sources of infection.
- Do not use the kits beyond shelf life.
- Do not mix different batches of reagents.
- ⚠ Warning:**
- To avoid error, do not forced to take out the cuvette from the device. Follow the device operation manual strictly, If the problem cannot be solved, contact the manufacturer for further technical support.

SYMBOLS USED ON LABELS

Symbol	Usage	Symbol	Usage
	Use-By date		Do not freeze
	Batch code		Biological risks
	Manufacturer		Do Not Reuse
	Temperature Limit		
	Contains sufficient for <n> tests		
	Do not use if package is damaged		
	Consult Instructions for use		
	Keep Away from Sunlight		
	In Vitro Diagnostic Medical device		
	Authorized Representative in the European Community		

References

Shang Hong, Wang Liu three, Shen Ziyu and so on. National Clinical Laboratory Practice (Fourth Edition) [M] Beijing: People's Medical Publishing House,2015:412-413.

Approval Date&Revision Date

- Approval Date: Sept 9,2015
- Revision Date: May 6,2016
- Revision Date: May 1, 2017
- Revision Date: Jan 1, 2021
- Revision Date: Apr 1, 2021
- Revision Date: Jan 1, 2023
- Revision Date: Dec 22,2023

尺寸:24*25cm展开尺寸,横向三折页再垂直方向两次对折