Instruction for C-Reactive Protein (CRP) QC

Product Name

C-Reactive Protein (CRP) QC

[Package Specification]

Applicable for PA50&PA54				
REF No.	Level	Specification		
32081021	QC(N)	0.2ml/bottle $ imes$ 2		
32081020	QC(H)	0.2ml/bottle $ imes$ 2		

Applicable for PA120					
REF No.	Level	Specification			
32081083	QC(N)	0.2ml/bottle $ imes$ 2			
32081082	QC(H)	0.2ml/bottle $ imes$ 2			

[Intended Use]

CRP QC is designed for quality control of C-Reactive Protein (CRP) detection by in vitro immune diagnostic system.

[Precaution]

This product is only used for in vitro diagnosis. Dispose the reagents according to the usual laboratory precautions.

Absorb the required amount for each use, and the remaining samples should not be returned to the original bottle.

If the quality control product is contaminated, the stability of the components will be decreased, it should be stop using and a new quality control product should be selected.

[Main Composition]

CRP antigen

Storage and Stability

The sealed detection kit can be stored at 2-8°C for 12 months. Do not freeze. Once opened, the reagents stored at 2-8°C are stable for 30 days.

[Instructions for use]

The quality control product is liquid ready-to-use type. The usage is as follows:

1. Remove bottles from the refrigerator(2-8 $^{\circ}$ C) and allow to warm to room temperature

for 5-10 minutes before mixing.

2. Gently invert at least 3 times before sampling, but avoid bubbles.

3. Open the bottle carefully to avoid ejecting the contents.

4. After sampling, cover the bottle cap as soon as possible and store it in a sealed condition of 2 ~ 8 $^\circ\!C$ in time.

For specific usage, please refer to the corresponding kit instruction.

[Assignment]

Assigned the quality control product in the company laboratory, using kit and specific protein analyzer, the methodology was Nephelometry.

The assignment result is shown on the QC bottle label.

[Icon Illustration]

LICOIL					
	\sim	Date of manufacture		Manufacturer	
	IVD	In vitro diagnostic medical device	4	Volume	
	Ŕ	Biological risks	REF	Catalogue number	

LOT	Batch code		Main component
	Use-by date	×.	Temperature limit
EC REP	Authorized representative in the European Community	CE	CE Marking

[Manufacturer]

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[Medical Device Manufacturing Enterprise Permit No.]

Guangdong SFDA (State Food and Drug Administration) authorized Medical Device Manufacturing Permit No. 20041046.

[Guarantee and Technical Support]

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