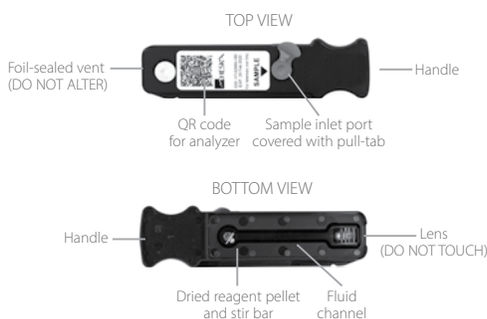


## Element i+ cCRP (Canine C-Reactive Protein) Cartridge Instructions

### Intended Use

The Element i+® cCRP (Canine C-Reactive Protein) assay is an in vitro diagnostic test for the quantitative determination of CRP in canine serum and lithium heparin plasma. This assay is used to detect and monitor systemic inflammation in canine patients. The reporting range of the assay is 0.4–20.0 mg/dL (4–200 mg/L). A result greater than 1.0 mg/dL (10 mg/L) is considered to be evidence of systemic inflammation.

### Element i+ Cartridge Description



### Principle of the Measurement

The Element i+ cCRP test uses a competitive immunoassay to generate a quantitative cCRP concentration output. When a specimen is added to a cartridge inlet port, it is mixed with a dried fluorophore-labeled cCRP. The mixture then reacts with anti-CRP immobilized on the cartridge sensor surface. cCRP in the sample competes with the fluorophore-labeled cCRP for binding to the anti-CRP antibody. Fluorescence illumination is by diode laser light coupled into the lens of the proprietary planar waveguide cartridge. Fluorescence imaging is used for signal transduction. The fluorescence generated is inversely proportional to the cCRP concentration of the specimen. Fluorescence intensity is converted to a quantitative cCRP concentration using cartridge lot-specific calibration information.

### Warnings

- ⚠ Do not touch the clear bottom of the cartridge.
- ⚠ Do not alter the silver foil seal on top of the cartridge. Do not use a cartridge with a punctured silver seal.
- ⚠ Do not use a cartridge dropped on the floor.

- ⚠ A new cartridge must be used for each measurement. The Element i+ Immunodiagnostic Analyzer will not allow a cartridge to be used more than one time.
- ⚠ Used cartridges should be disposed of as biohazard waste in accordance with local laws.

### Additional Equipment

- Element i+ Immunodiagnostic Analyzer
- 100 microliter fixed volume mini-pipette (supplied with analyzer and available separately)
- Pipette tips (supplied with analyzer and available separately)

### Specimen Requirements

The test has been designed to be run with fresh canine serum or lithium heparin plasma samples. Other plasma sample types have not been tested.

Avoid using plasma or serum with precipitates. Do not dilute the sample.

### Running a Test

1. Obtain a serum or lithium heparinized plasma sample.
2. On the main screen of the analyzer, touch Worklist or Manual Test.
3. In Worklist Mode, confirm that all fields have correct information and then touch ✓ to proceed.
  - In Manual Mode, enter sample information in the required fields.
  - Touch ✓ to proceed.
4. Open the pouch by tearing at the notch. Carefully remove the cartridge by the handle and place it on a flat surface.
 

NOTE: Do not touch the bottom of the cartridge.

NOTE: If the cartridge was refrigerated, allow to warm to room temperature for at least 15 minutes before opening the pouch.

NOTE: Cartridge must be used within 1 hour of removal from pouch.
5. With the cartridge flat on the bench, remove the pull-tab from the sample inlet port and discard. Use the handle to steady the cartridge while removing the pull-tab.
 

NOTE: Cartridge must be used within 15 minutes of removing pull-tab.

6. Affix a tip to the 100 µL fixed volume mini-pipette. Aspirate 100 µL of sample, insert the pipette tip into the inlet port hole, and dispense the full sample amount into the hole.
  7. Touch on the Prepare Sample screen to open the analyzer door.
    - Insert the cartridge until you feel a click and hear a beep.
  8. The test will run automatically. A status bar and countdown timer will display on the screen and the indicator light on the front of the analyzer will blink to indicate a test is running: To cancel during the run, touch X at the upper right of the screen.
  9. Upon test completion, patient results will display on the screen.
    - Touch (home button) to exit results screen.
    - The screen will provide an indication when it is safe to remove the used cartridge.
- NOTE: Do not attempt to remove cartridge before signaled by analyzer.

## Canine Reference Interval

0.0–1.0 mg/dL (0–10 mg/L)

## Performance Characteristics

Reporting range:

0.4 to 20.0 mg/dL (4–200 mg/L)

Reference method:

Gentian cobas canine CRP Immunoassay

Accuracy:

Overall bias based on the Passing-Bablok slope is less than 20% within the 95% confidence interval for samples  $\geq 1.0$  mg/dL (10 mg/L) cCRP.

Precision:

Coefficient of Variation  $\leq 15\%$  for samples  $\geq 1.0$  mg/dL (10 mg/L) cCRP.

## Known Interfering Substances

Interference	Interfering Substance and Conditions Tested	cCRP Sample Concentrations Tested	Result
Hemolysis	Hemoglobin up to 500 mg/dL	1.0 and 7.5 mg/dL (10 and 75 mg/L)	No significant effect
Icterus	Bilirubin up to 20 mg/dL		
Lipemia	Lipids/Triglycerides up to 1000 mg/dL		

## Storage and Shelf Life

Storage:

35.6–77°F (2–25°C)

Expiration date:

Printed on the cartridge pouch.



For further assistance, please call Heska's Medical and Technical Consultants at

US 800 464 3752  
www.heska.com

CA 866 382 6937  
www.heskavet.ca

AU 1300 437 522  
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