# (EN) CueSee® VeriSTAT QC - Level 1

vitro diagnostic medical devices and carries the CE mark.

POCT analyzer. Each ampule contains 2.5 mL of solution.

CueSee® VeriSTAT QC - Level 1 is an assayed aqueous quality control material for

professional use in the verification of the precision and accuracy of the Abbott i-STAT®

It is intended that CueSee® VeriSTAT QC - Level 1 should be used in the periodic

verification of the precision, accuracy of the Abbott i-STAT POCT analyzer when

measuring: pH, pO2, pCO2, Na+, K+, Ca++, Cl-, Glucose, Lactate, Urea, Creatinine,

CueSee® VeriSTAT QC - Level 1 complies with the European Directive 98/79/EC on in

CueSee® VeriSTAT QC - Level 1 complies with the following US codes of Federal

CueSee® VeriSTAT QC - Level 1 is an assayed aqueous quality control material for

professional use in the verification of the precision and accuracy of the Abbott i-STAT®

CueSee® VeriSTAT QC - Level 1 quality controls are prepared using salts in an

aqueous, physiologically buffered matrix. Tonometry with predetermined levels of

oxygen and carbon dioxide balanced with nitrogen and different salt concentrations

provides distinct levels for each parameter, simulating clinically significant ranges

of acid-base and electrolyte balance, respiratory function, glucose, lactate,

urea and creatinine concentrations, within the reportable range of the Abbott

CueSee® VeriSTAT QC - Level 1 must be stored at a temperature of 2-8  $^{\circ}$ C (35–46  $^{\circ}$ F). Stored unopened at this temperature it is guaranteed stable until the expiration date as indicated on the ampule and on the outer box. After opening the CueSee® VeriSTAT QC - Level 1 ampule, the product is stable for 30 seconds.

1. Take the ampule out of its packaging and equilibrate the ampule to room temperature (20-25 °C/68-77 °F) for a minimum of 1 hour, but not more than 10 days. Note: Do not

2. Immediately before use, shake the ampule vigorously for at least fifteen seconds to

3. Swirl the ampule gently to return the solution to the bottom of the ampule. Allow any

6. Using one of the techniques described under "Instructions for Blood Gas/Electrolyte/

7. Refer to CLIA regulations, state regulations, accrediting agency requirements as well

Metabolite Cartridges" in the Quality Control section of the i-STAT system manual,

transfer sample from the ampule to the test cartridges within 30 seconds of opening

re-equilibrate the gases with the solution by holding the ampule between the thumb

put ampules back into refrigerator once exposed to room temperature.

5. The One Point Cut (OPC) ampule must be opened as presented below.

Regulations (CFR): 42 CFR part 72 and 21 CFR parts 606, 640 and 820.

CueSee® VeriSTAT QC - Level 1 is for professional in vitro diagnostic use only.

Intended Use

POCT analyzer.

TCO2 and Hematocrit.

**IVD Medical Device** 

Summary

Reagents

i-STAT POCT analyzer.

Storage and stability

and index finger.

the ampule.

bubbles to rise before opening the ampule. 4. Protect fingers with gauze, tissue or gloves.

as the i-STAT system manual for frequency of use.

Procedures

#### Precautions

## 1. CueSee® VeriSTAT QC - Level 1 is for use on the Abbott i-STAT analyzer only.

- 2. For in vitro diagnostic use only.
- 4. CueSee® VeriSTAT QC Level 1 is not to be used as a calibrator.
- 3. Consult local environmental authorities for proper disposal.

#### **Reference Values**

Enclosed values have been obtained by equilibrating randomly selected ampules from the applicable batch at 25 ± 1 °C (77 ± 2 °F) before measurement. The values have been obtained on multiple i-STAT instruments using multiple cartridge types.

The value ranges indicate the values for each parameter within which the obtained results must fall.

## Please Note

- The values in the table are applicable at sea level for the Abbott i-STAT POCT analyzer with the applicable cartridge types and cartridge lots only.
- Due to variations in CLEW updates, cartridge type and cartridge lot number, for some analyte concentrations in CueSee® VeriSTAT QC Level 1, results may fall outside of the i-STAT reportable range. If necessary, it is recommended that the Calibration Verification pathway be used to obtain a numerical result.
- Incorrect sampling, storage, or other mishandling may cause the readings to deviate from the target values.
- The pO2 values of CueSee® VeriSTAT QC Level 1 controls vary inversely with temperature changes. To obtain a high degree of correlation with the values in the table, the ampules should be equilibrated as close to 25 °C (77 °F) as possible.
- CueSee® VeriSTAT QC Level 1 controls are very sensitive to room air contamination. To minimize the effect of air contamination, sample within 30 seconds of opening. The effect of room air on pH is negligible.

This product has been manufactured according to Eurotrol specifications.

For further information please contact:

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- T: 1-978-598-3779
- www.eurotrol.com

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LOT

r0/

REF

Batch code

CE mark

Temperature limitation

Reference number

Eurotrol B.V., Keplerlaan 20, 6716 BS Ede, The Netherlands

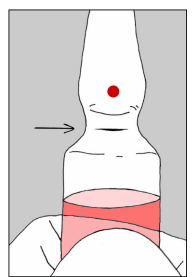
Symbols used

F: 1-978-598-3780

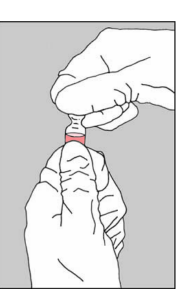
- Attention, see instructions for use
- Use by
- IN Vitro Diagnostic Medical Device
- Manufacturer

i-STAT is a registered trademark of Abbott Laboratories

Hold the bottom of the ampule with thumb pointing to the red colored dot.



Cut located below the red dot is the breaking point of the ampule.



Grasp the top of the ampule with other hand, positioning thumb at the red dot.



Press back to break at the cut under the red dot.

