Nova Biomedical STATSensor - Creatinine (LTR44807)



Approval Workgroup: POCT

Folder Name: POCT\03 POCT-Waived Test

Procedures

Revision: 2.00 Next Review Date:

Last Approved By: Benirschke, Robert (7/7/2016

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Background Information	These are the instructions for the creatinine test using the Nova Biomedical Stat Sensor.
Intended Use	This method is used to perform quantitative measurements of creatinine in whole blood.
Manufacturer	This method is manufactured by Nova Biomedical. It is performed as per the manufacturer's instructions, without any in-house modifications.
OSHS Safety Classification	This procedure is classified as a Category I potential hazard by OSHA standards. Appropriate barrier equipment and precautions must be used in performing the procedure.
CLIA Complexity Level	Waived
Summary and Explanation of Test	Creatinine is breakdown product of creatine phosphate in muscle tissue. Creatinine measurements are used in the diagnosis and treatment of renal diseases, in monitoring renal dialysis, and as a calculation basis for measuring other urine analytes.
Principle	Creatinine is measured in whole blood by an electrochemical method.
Specimen Requirements	Whole blood collected from a fingerstick.
Materials and Equipment Needed	Nova Biomedical Stat Sensor Non sterile gloves, lancet and alcohol prep pad, gauze Nova Biomedical Stat Sensor test strip Nova Biomedical Creatinine controls, levels 1, 2, 3.
Reagent Preparation and Storage Instructions	 Test strips and control material is stored at a refrigerated temperature of 2 to 8°C. Reagent solutions once opened and refrigerated are stable for 90 days or until the expiration date which occurs first. Test strip vials once opened are stable for 90 days. Keep test strips tightly sealed in the storage container. Sealed test strips are stable until the expiration noted on the container. Discard all test strips and solutions once past their expiration date. NOTE: Test strips and QC material MUST be brought to

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4. Check the strip lot number and press **Accept**.



5. Scan the patient wristband, the patient identifier is the CSN. The CSN can be manually entered, however, double check the entry before acceptance to insure results post on the correct EPIC record.



6. Insert a test strip into meter.



- 7. Wash the patient's hand thoroughly and massage the finger to stimulate blood flow.
- 8. Use a safety lancet to puncture the finger. Gently squeeze the finger to form a blood drop.
- 9. Touch the test strip to the blood drop. The creatinine result will appear on the screen in 30 seconds.



Note: the test strip must fill completely after touching the blood droplet. If the test strip does not fill completely, do not touch the blood droplet a second time. Discard the test strip and repeat the test with a new test strip.

10. To accept the result, press **Accept**. To reject the result, press **Reject**. A comment can be appended to the result, press Comment on the right hand bottom of the result screen.



11. To review other test results, press **Review** from the Patient Test screen.



Calculations None Quality Control Quality control must be performed every 24 hours to insure accuracy and precise of the device. 1. Follow steps 1 and 2 in the procedure above. 2. From the Patient Test screen, press QC.



- 3. Enter the strip lot number as described above in the procedure and press **Accept**.
- 4. Enter the QC lot number and press **Accept**.
- 5. Insert the Test Strip as described above in the procedure.
- 6. Touch a drop from the QC bottle to the test strip.



- 7. Test strip must fill completely upon touching the QC droplet. **DO NOT** add a second QC drop to the test strip. Discard the test strip and insert a new test strip into meter.
- 8. Result will appear in 30 seconds



9. To accept the result press **Accept**.

NOTE: Patient results cannot be reported unless all three levels of QC are within the acceptable range.

Result Reporting

The Stat Sensor Creatinine meter must be placed in the docking station after to facilitate the reporting of the results. The system is interfaced to Nova Net and Telcor and results will automatically flow into EPIC. The instrument sends across to NovaNet and Telcor the result, date, time

Fei Ma	rum: male: 0.5 - 1.2 mg/dL le: 0.7 - 1.4 mg/dL truments used for Point of Care testing that are used on more than
Cleaning the Motor	
one dis	e patient must be disinfected with an approved hospital grade infectant after use per Infection control Policy 10.7 Non Dedicated item Care Equipment.
	Acceptable creatinine range established by Radiology is 0.8 - 1.5 mg/dL. Results falling outside of Radiology established range of 0.8 - 1.5 mg/dL will need to be confirmed utilizing the laboratory and notifying the ordering physician. Acceptable eGFR value established by Radiology is >30ml/min for MRI procedures and >60ml/min for CT procedures.
4.	GFR results of <30ml/min for MRI procedures and <60ml/min will need to be confirmed utilizing the laboratory and notifying the ordering physician. Refer to Contrast Media Administration CP08-3732 and Gadolinium Based Contrast media Administration CP08-3721
5.	The Nova Stat Sensor has a reportable range of 0-8 mg/dL.
6.	If meter is dropped, DO NOT use for patient testing. Call POC supervisor for assistance. QC and linearity testing will be run to assure there is no instrument failure.
7.	If strip appears irregular in any way do not use for testing.
8.	If POC creatinine testing is not available due to lack of supplies or analyzer problems, send appropriate sample to lab for testing
9.	If meter is not working please call POC supervisor at EV (772-4684) or pager 8585 for assistance during regular business hours. If problem occurs outside of regular business hours someone will contact you the next business day.
Limitations Wh	nole blood is the only acceptable specimen for analysis
in I	ne Stat Sensor is not working properly, order a STAT Creatinine level EPIC and send a venipuncture collected specimen to the to the poratory for analysis.
References 1. 2.	Nova Biomedical Stat Sensor Procedure Manual. Clinical Practice Radiology Manual, NorthShore University HealthSystem, April 2010.
Effective Date The	e effective date for this procedure is June 30, 2010.
Written by Ro	bert Rosecrans, Ph.D.