



PROSTATE CANCER **BIOREPOSITORY NETWORK**

SOP No: 005  
QC of DNA

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| <b>STANDARD OPERATING<br/>PROCEDURE</b>           | SOP No. 005<br>QC of DNA                               |
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# PROSTATE CANCER **BIOREPOSITORY NETWORK**

SOP No: 005  
QC of DNA

## 1. PURPOSE

To describe the procedure for the QC of DNA extracted from frozen tissue by real-time PCR. NOTE: This SOP does not cover detailed procedures for handling Human Biological Materials or hazardous chemicals and it is recommended that personnel following this SOP refer to institutional safety guidelines.

**The following SOP for QC of DNA is followed by the JHU, NYU and UW Network Sites of the PCBN.**

## 2. RESPONSIBILITIES

**Authorized personnel** performing QC of DNA extracted from frozen prostate tissue must ensure that:

- all procedures are followed correctly
- all samples are adequately coded during processing
- all documentation is completed, and accurate records maintained on all samples

## 3. HEALTH AND SAFETY

Personnel carrying out this procedure must maintain safe working practices and observe all relevant Environmental Health and Safety (EH&S) guidelines. This includes the appropriate use of Personal Protective Equipment (PPE), and the procedures for waste disposal, disinfection & spill clean-up and biosafety.

## 4. EQUIPMENT AND MATERIALS FOR JHU & NYU NETWORK SITES

| Equipment                                  | Materials   |
|--|---|
| iCycler real time Thermocycler<br>Nanodrop | 96-well plates<br>Primers as described<br>SYBR Green PCR Master Mix<br>(Invitrogen Catalog # 4309155) |

## 5. EQUIPMENT AND MATERIALS FOR UW NETWORK SITE

| Equipment   | Materials  |
|---|--|
| Gel electrophoresis system<br>Gel Doc<br>Nanodrop | Agarose<br>TAE Buffer<br>6X Sample Loading Buffer<br>DNA Ladder Standard |



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## 6. PROCEDURES FOR JHU AND NYU NETWORK SITES ONLY

- Quantitate DNA samples extracted using **PCBN SOP004 DNA Extraction from Frozen Tissue** by Nanodrop. Refer to the Nanodrop user guide for instructions of use.
- If DNA concentration is greater than 100 ng/ul, dilute DNA sample to 100 ng/ul.
- QC of DNA samples extracted using **PCBN SOP004 DNA Extraction from Frozen Tissue** by real-time PCR.

- Rxn mix (20 µl): µl/rxn

|            |    |
|------------|----|
| gDNA       | 2  |
| SYBR Green | 10 |
| F-primer   | 1  |
| R-primer   | 1  |
| dH2O       | 6  |

- A.  $\beta$ -globin primers (100 bp product)

Forward: 5'-GTGCACCTGACTCCTGAGGAGA-3'  
Reverse: 5'-CCTTGATACCAACCTGCCAG-3'

- B. 18S primers (110 bp product)

Forward: 5'-GATGGTAGTCGCCGTGCC-3'  
Reverse: 5'-GCCTGCTGCCTTCCTTGG-3'

- Cycling parameters

Cycle 1: ( 1X)  
Step 1: 95.0°C for 03:00  
Cycle 2: ( 35X)  
Step 1: 95.0°C for 00:30  
Step 2: 60.0°C for 00:30  
Step 3: 72.0°C for 00:45

## 7. PROCEDURES FOR UW NETWORK SITE ONLY

- Quantitate DNA samples extracted using **PCBN SOP004 DNA Extraction from Frozen Tissue** by Nanodrop. Refer to the Nanodrop user guide for instructions of use.
- Measure the  $A_{260}/A_{280}$  ratio for an estimate of DNA purity with respect to contaminants that absorb UV.
- DNA length can be determined using agarose gel electrophoresis.
- DNA samples with sharp 10kb band are considered as high-quality DNA.