

BGI Sample Testing Report

1. Project Information

Report No.: THKa15093002

Project Name	UW - Sam **** - Sam White - de-novo genome Panopea generosa	Project No.	F15FTSUSAT0328
Customer Name	Steven Roberts	Customer Unit	Univ. of Washington
Lab Sample Collector	lam tsz tung	Lab Sample Receiving Date	20150929
Lab Sample Tester	Lam Tsz Tung	Lab Sample Testing Date	20150929
Reported by	YUEN Ka Yiu Inspected lamtsztung	Approved by	Wong Wai Man Report 20150930 Vivien Date

2. Sample Test Method

①Method of concentration determination: ■Qubit Fluorometer、□NanoDrop、□Microplate Reader

②Method of sample integrity test: ■Agarose Gel Electrophoresis

3. Sample Test Result

No.	Sample Name	Sample Number	Tube No.	Concen- tration(ng/ μL)	Volume(μL)	Total Mass(µg)	sample Integrity	Library type	Test result	Remark
1	gDNA geoduck	8521509003287	1	22. 8	790	18. 01	Degraded slightly	≤800bp Insert Size	Level A	

Note*:

- 1. The test result based on the $\langle DNA \rangle$ sequencing sample quality standards explains whether the testing sample meets the requirement of library construction.
- a) Level A means the sample is qualified, and the amount of sample is sufficient for two or more library constructions.
- b) Level B means the sample is qualified, but the amount of sample only satisfies one time library construction.
- c) Level C means the sample does not totally meet the requirements of library construction and sequencing. BGI can try to construct the library but the quality of the sequence is not guaranteed
- d) Level D means the sample does not meet the requirements of library construction and sequencing. BGI does not suggest in using this sample.
- 2. According to BGI's data, the one-time successful rate of library construction is more than 95% for samples of level A and level B.
- 3. According to BGI's data, the risks of library construction for sample of level C or level D are listed below:
- a) The deficiency of the quantity: There may be the risk of failure in library construction and the yield of library of experiment may be too low to sequencing, and the database of low yield for sequencing may lead to poor randomness.
- b) Degradation of sample: It may cause high duplication rate of library and insert fragment will be abnormal."
- c) Pollution by Protein or Insoluble Impurity: It may affect the fragmentation effect, leads to insert size unstable, influence the SNP analysis
- d) RNA contamination: It possibly effects the DNA concentration quantitative accuracy.
- 4. If the partner insists on using the sample of level C or level D, the risk and responsibility is taken by the cooperative partner.
- 5. Other notes:
- A) Sample is contaminated by RNA. RNase A treatment is recommended.

4. Appendix

Appendix 1: Test results of Qubit Fluorometer or Microplate Reader

Appendix 2: Test results of Agarose Gel Electrophoresis

Appendix 3: Original information of sample

5. Statement

1. The results shown in this report refer only to the sample of the report unless otherwise stated.

2. This test report cannot be copied partly without the prior written permission of the Lab.

Appendix 1: Test results of Qubit Fluorometer or Microplate Reader

1. Pre-treatment

After the sample melted the ice, centrifuged and fully mixed, take appropriate samples for testing.

2. Test Result

Sample Name	Sample Number	Test Instrument	Test Kit	Dilution Ratio(×)	Test Volume (μL)	Test Concentration(ng/ µL)	Concentration of original sample(ng/µL)	Remark
gDNA geoduck	8521509003287	Qubit	DNA BR	1	1	22. 8	22.8	

Appendix 2: Test results of agarose gel electrophoresis

1. Pre-treatment

After the sample melted the ice, centrifuged and fully mixed, take appropriate samples for testing.

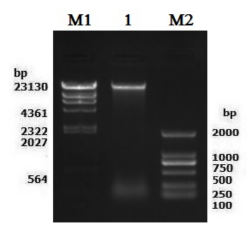
2. Test Parameter

Concentration of Agarose Gel: 1 %; Voltage: 150 V; Electrophoresis

Time: 40 min

3. Test Result

(1) Electrophoretogram:



Lane No.	Sample Name	Dilution Ratio(\times)	Test Volume(μL)	Sample Integrity	Remark
M1	λ-Hind III digest(Takara)	1	3		
1	gDNA geoduck	1	4. 39	Degraded slightly	RNA contamination
M2	D2000 (Tiangen)	1	6		

Appendix 3: Original information of sample

Sample Type:									
Genome DNA									
Sample sta	Sample status:								
Dissolved in	n 10mM Tris-	HC1							
Further In									
Sample Name Species No. of Concentration(ng/Volume(µL) Total Fragment Size OD260/280 OD260/230 Remark									
gDNA geoduck	Panopea generosa	1	168. 00	500.00	84		1.84	0.87	