DNBseq[™] service overview RNA-Seq (Transcriptome) Sequencing

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Service Description

Transcriptome sequencing is used to reveal the presence, quantity and structure of RNA in a biological sample under specific conditions. Compared to hybridization-based RNA quantification methods such as microarray analysis, sequencing-based transcriptome detection can quantify gene expression with low background, high accuracy and high levels of reproducibility within a large dynamic range. In addition, transcriptome sequencing does not require an existing genome sequence and can detect mutations, splice variants and fusion genes that cannot be detected by microarrays.

Sequencing Service Specification

BGI transcriptome sequencing services are executed with the DNBseq sequencing technology, featuring cPAS and DNA Nanoballs (DNB^m) technology for superior data quality.



Sample Preparation and Services

- \cdot Multiple choices for mRNA enrichment and rRNA removal kits
- \cdot Stranded and non-stranded sequencing is available
- \cdot 100bp and 150bp paired-end sequencing options available
- $\cdot \geq$ 30 Million reads per sample recommended
- Raw data and bioinformatics analysis are available in standard file formats
- · Advanced and custom bioinformatics data analysis
- \cdot Cloud-based data storage and delivery system



Sequencing Quality Standard

 \cdot Guaranteed ${\geq}80\%$ of bases with quality score of ${\geq}Q30$



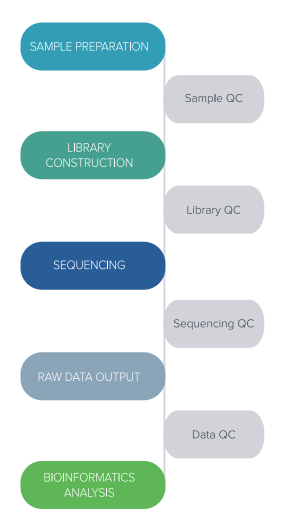
Turnaround Time

- Typical 30 working days from sample QC acceptance to filtered raw data availability
- Expedited service are available, contact your local BGI specialist for details



Project Workflow

We care for your samples from the start to the result reporting. Highly experienced laboratory professionals follow strict quality procedures to ensure the integrity of your results.





DNBseq Sequencing System

DNBseq is an innovative high-throughput sequencing solution, developed by BGI's Complete Genomics subsidiary in Silicon Valley. The system is powered by combinatorial Probe-Anchor Synthesis (cPAS), linear isothermal Rolling-Circle Replication and DNA Nanoballs (DNB[™]) technology, followed by high-resolution digital imaging.



The combination of linear amplification and DNB technology reduces the error rate while enhancing the signal. The size of the DNB is controlled in such a way that only one DNB is bound per active site on the DNBseq flow cell. This densely patterned array technology provides optimal sequencing accuracy and increases flow cell utilization.

Data Analysis

In addition to raw data output, BGI offers a range of standard and customized bioinformatics pipelines for your transcriptome sequencing project. Reports and output data files are delivered in industry standard file formats: FASTQ, BAM and Excel.

Standard Analysis

- Quantitative expression profiles
- Alternative splicing analysis
- Fusion gene analysis
- Time series analysis
- Gene ontology analysis

- Pathway enrichment analysis
- Hierarchical clustering analysis
- Protein-Protein Interaction (PPI) analysis
- Fungal pathogenic gene annotation (for fungi)
- Plant disease resistance genes annotation (for plant)

Customized Analysis

Further customization of Bioinformatics analysis to suit your unique project isavailable: Please contact your BGI technical representative.

Sample Requirements

We can process your total RNA, blood, cell line, FFPE, fresh frozen tissues and single cell samples from a variety of species, with the following general requirements:

Sample Type	Species	Amount	Concentration (ng/µL)	RIN/RQN Value	28S/18S	DV ₂₀₀
Total RNA	Human	≥200ng	≥20	≥7.0	≥1.0	N/A
	Human (FFPE)	≥200ng	≥70	N/A	N/A	≥30%
	Mouse/Rat	≥200ng	≥20	≥7.0	≥1.0	N/A
	Insect	≥1µg	≥40	N/A	N/A	N/A
	Other Animals	≥1µg	≥40	≥7.0	≥1.0	N/A
	Plant /Fungi	≥1µg	≥40	≥6.5	≥1.0	N/A

Sample Type (For human samples)	FFPE	Whole Blood	Cell Line	Tissue
Requirement	≥5 slides ≥5 µm slice per slide	≥1mL	≥2X10⁵ cells	≥30mg

Low-input transcriptome sequencing is available.

Stable and High-Quality Data performance

1,072 samples were randomly selected from over 10,000 samples that were sequenced at BGI's laboratory over a period of 6 months. The data output and data quality remained stable over that period. The average Q20 and Q30 scores were 97% and 89.5% respectively.



Stable data quality scores from DNBseq production line



Request Information or Quotation

Contact your BGI account representative for the most affordable rates in the industry, to discuss how we can meet your specific project requirements or for expert advice on experiment design, from sample to bioinformatics.

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For BGI's scientific publications relating to DNBseq Small RNA Sequencing, sample shipping instructions or sample submission forms, please visit our website.

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All Services and Solutions are for research use only.

DNBseq Services are executed in our service laboratory in China.



We Sequence, You Discover