RNA-Seq Technical Specifications

GENEWIZ[®] From Azenta Life Sciences

RNA Sequencing Services

- Standard RNA-Seq
- Strand-Specific RNA-Seq
- Small RNA-Seq
- Ultra-Low Input RNA-Seq

- Single-Cell RNA-Seq*
- Long-Read RNA-Seq using PacBio or Oxford Nanopore Technologies*

Sample Preparation

Our extraction solutions are compatible with over 30 standard and

hundreds of custom sample types to accommodate your starting material. The below table is by no means an exhaustive list.

Minimum Amount[†]

500 ng (standard)

10 pg (ultra-low)

1 cell (ultra-low)

10⁶ cells

2 mg

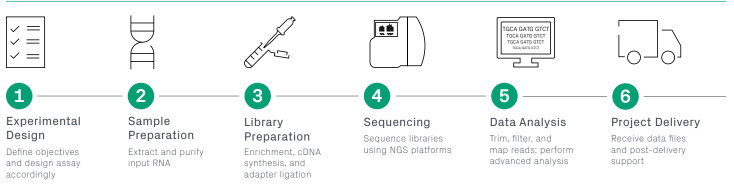
2 slides

10⁴ cells (standard)

- Digital Spatial Profiling*
- CLIA RNA-Seq*

*Not covered here. See genewiz.com for more details.

RNA Sequencing Workflow



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Sample Type*

Eukaryotic cell

Prokaryotic cell

Frozen tissue

Total RNA[‡]

pellet

pellet

FFPE

1 Experimental Design

We offer resources to help you find the best NGS solution and experimental design for your project.



Interactive NGS Solution Selection Tool: genewiz.com/ngs



Contact us for a **free technical consultation** with a Ph.D.-level scientist

Other sample types accepted. View <u>Sample Submission Guidelines</u> for details. Please inquire about submitting lower inputs.

¹Contact us about our RNA Stabilization Tubes to ship RNA samples at ambient temperature.



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Recommended

Amount

10⁶ cells

10⁸ cells

10 mg

4 slides

2 µg

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Library Preparation

RNA-Seq Service	Target RNA	RNA Selection Method
Standard & Strand- Specific	mRNA (eukaryotic)	Poly(A) selection
	mRNA + IncRNA	rRNA depletion
Small	Small RNA (miRNA, siRNA, piRNA)	Size fractionation with adapter ligation to 5' phosphate
Ultra-Low Input	mRNA (eukaryotic)	Poly(A) selection with enrichment for full- length transcripts

Sequencing

Platform	Illumina® NovaSeq™	
Configuration	2×150 bp	
Depth	Customizable to your project needs*	
Data Quality	Guaranteed ≥80% bases with Q30 or higher	

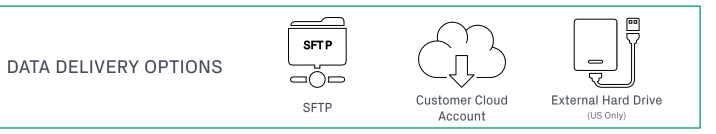
*Generally, we recommend 5-10 million read pairs per sample for small genomes (e.g. bacteria) and 20-30 million read pairs per sample for large genomes (e.g. human, mouse). Medium genomes often depend on the project, but 15-20 million read pairs per sample is typically sufficient. For *de novo* transcriptome assembly projects, we recommend 100 million read pairs per sample.

5 Data Analysis

RNA-Seq Service	Standard Analysis Package	Additional Analysis Options
Standard Strand-Specific Ultra-Low Input	 Trimming Mapping Differential gene expression 	 Gene fusion discovery RNA SNP/INDEL detection Novel transcript discovery <i>De novo</i> transcriptome assembly
Small	 Trimming Mapping Differential gene expression Small RNA discovery 	

6 Project Delivery

Deliverables for All Projects	Optional Deliverables
• Sample quality control report • Raw data (FASTQ files)	 Aligned data (BAM file) Hit counts (TXT file) DGE results (CSV file) GO enrichment analysis (CSV file) Differential splicing analysis (DEXSeq report) De-multiplexed, aggregated Picard BAM file with summary metrics





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