

## **Eukaryotic RNA Sequencing**

#### 1.Sample Requirements

Library Type	Sample Type	Amount	Volume	Concentration	RNA Integrity Number (Agilent 2100 <sup>™</sup> )	Purity (NanoDrop <sup>™</sup> )
Eukaryotic RNA- Seq (cDNA library)	Total RNA	≥ 200 ng	≥ 10 µL	≥ 20 ng/μL	≥ 4.0, with smooth base line	OD260/280 ≥ 2.0; OD260/230 ≥ 2.0; No degradation, No contamination
	Total RNA (Blood)	≥ 400 ng	≥ 20 μL	$\geqslant$ 20 $\mu$ L	≥ 5.8, with smooth base line	
	Total RNA (Single Cell)	≥ 100 ng	≥ 20 µL	≥ 10 ng/µL	≥ 5.8, with smooth base line	
	Amplified cDNA (double- stranded)	≥ 100 ng	≥ 10 µL	≥ 10 ng/µL	Fragments should be distributed between 400bp - 5000bp with main peak at ~2000 bp	OD260/280 ≥ 2.0 OD260/230 ≥ 2.0; No degradation, no contamination
Eukaryotic RNA- Seq (strand specific library)	Total RNA	≥ 400 ng	≥ 20 μL	≥ 20 ng/µL	≥ 5.8, with smooth base line, with smooth base line	OD260/280 ≥ 2.0; OD260/230 ≥ 2.0; No degradation, no contamination

For total RNA less than 100 ng, please contact us for ultra-low input solutions.

### 2.Sequencing Parameters

Platform	Illumina NovaSeq 6000			
Read length	Paired-end 150			
Recommended sequencing depth	≥ 20 million read pair per sample for species with reference genome;> 50 million read pairs per sample for species without reference genome (de novo transcriptome assembly projects)			
Data quality	Guaranteed ≥ 85% bases with Q30 or higher			
Turnaround time	Within 2~3 working weeks from library construction verification to data releasing without bioinformatic analysis. (depending on the sample size);			

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### 3.Data Analysis Contents

Standard analysis					
Data filtering					
Transcriptome assembly & Gene functional annotation (only for species without reference genome)					
Mapping to reference genome/assembled genome					
Gene expression quantification & Differential expressed genes profiling & Enrichment analysis					
Protein-Protein Interaction (PPI) analysis					
Transcription factors functional annotation analysis					
Oncogene functional annotation analysis					
SNP & InDel analysis					
Alternative splicing analysis					
Fusion gene prediction (Only for tumor sample and cancer cell line)					