

ChIP-Seq

1. Sample Requirements

Sample Type	Amount	Volume	Concentration	Fragment size	Purity
Enriched DNA Sample	≥ 10 ng	\geqslant 20 μ L	≥0.5ng/μL	Main peak should be 100 bp~500 bp.	OD260/280=1.8-2.0 no degradation, no contamination

2. Sequencing Parameters

Platform	Illumina NovaSeq 6000			
Read length	Paired-end 150 bp			
Recommended sequencing depth	\geq 20 million read pairs per sample for the species with reference genome			
Data quality	Guaranteed ≥ 85 % bases with Q30 or higher			
Turnaround time	Typical 18 wording days for 24 or fewer samples from project verification to data releasing			

3. Data Analysis Contents

Standard analysis

Data quality control

(get rid of reads containing adapter or with low quality; Q20, Q30, error rate distribution, GC distribution, total bases)

Mapping onto reference genome (mapping rate, reads distribution)

Peak calling

Motif prediction

Peak annotation (downstream or overlapping gene, TSS) and functional analysis of peak-associated genes (GO, KEGG)

Summary of differential peaks and functional analysis of differential peak related genes

Visualization of ChIP-seq data

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