

# **Whole Genome Bisulfite Sequencing** (Gene Methylation)

### 1. Sample Requirements

| Sample Type | Required Amount | Volume  | Concentration | Purity   |
|-------------|-----------------|---------|---------------|--|
| Genomic DNA | ≥ 200ng         | ≥ 20 μL | ≥ 10 ng/μL    | OD260/280=1.8-2.0;<br>0 < OD260/230 < 3;<br>No degradation or<br>contamination |

#### 2. Sequencing Parameters

| Platform                        | Illumina NovaSeq 6000  |  |  |
|---------------------------------|--|--|--|
| Read length                     | Paired-end 150   |  |  |
| Recommended<br>Sequencing Depth | $\geq$ 30 $\times$ coverage for the species with reference genome;   |  |  |
| Data quality                    | Guaranteed ≥ 80% bases with Q30 or higher  |  |  |
| Turnaround time                 | Typical 6~7 weeks for fewer than 20 samples from project verification to data releasing without bioinformatic analysis |  |  |

## 3. Data Analysis Contents

#### **Standard analysis**

Data quality control (filtering reads containing adapter or with low quality; Q20, Q30, error rate distribution, GC distribution, total bases)

Mapping onto reference genome (mapping rate, duplication rate, sequencing depth, reads coverage)

mCs detection, methylation level calculation

- (1) Methylation level and frequency distribution in different sequence context (CG, CHG, CHH)
  (2) Methylation level and frequency distribution in different chromosomes
  (3) Methylation level and frequency distribution in different functional elements (promoter, 5'UTR, exon, intron, 3'UTR)

Differentially Methylated Site (DMS) detection

Differentially methylated regions (DMRs), Differentially Methylated Promoter (DMPs) detection and annotation

Function enrichment (Gene Ontology and KEGG Pathway) of DMR-associated genes and DMP-associated genes

Visualization of BS seq data