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# **GSA Submission**

## \*Alias

Submission name of the GSA. This field is used when the record does not yet have an

accession and needs to be referenced by other objects.

## \*Data Released

Select Release on specified dateor give release data in correct format (yyyy-MM-dd).

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# Experiments

## **Meta Information**

## \*Platfom

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Thesequencing platform and instrument model

Instrument Model
454 GS 20
454 GS FLX
454 GS FLX Titanium
454 GS FLX+
454 GS Junior
AB 310 Genetic Analyzer
AB 3130 Genetic Analyzer
AB 3130xL Genetic Analyzer
AB 3500 Genetic Analyzer
AB 3500xL Genetic Analyzer
AB 3730 Genetic Analyzer
AB 3730xL Genetic Analyzer
AB 5500 Genetic Analyzer
AB 5500xl Genetic Analyzer
AB SOLiD 3 Plus System

AB SOLiD 4 System
AB SOLiD 4hq System
AB SOLID PI System
AB SOLiD System 1.0
AB SOLiD System 2.0
AB SOLiD System 3.0
Complete Genomics
HelicosHeliScope
Illumina Genome Analyzer
Illumina Genome Analyzer II
Illumina Genome Analyzer IIx
IlluminaHiScanSQ
IlluminaHiSeq 1000
IlluminaHiSeq 1500
IlluminaHiSeq 2000
IlluminaHiSeq 2500
IlluminaMiSeq
Ion Torrent PGM
Ion Torrent Proton
PacBio RS
IlluminaNextseq 500
IlluminaHiSeq 3000
IlluminaHiSeq 4000
IlluminaHiSeq X-10
Agilent Infinity 1290 UHPLC - 6550 QTOF MS
454 GS 20
454 GS FLX
454 GS FLX Titanium
454 GS FLX+

454 GS Junior
AB 310 Genetic Analyzer
AB 3130 Genetic Analyzer
AB 3130xL Genetic Analyzer
AB 3500 Genetic Analyzer
AB 3500xL Genetic Analyzer

## \*Alias

Submission name of the experiment. This field is used when the record does not yet have an accession and needs to be referenced by other objects.

## \*Title

Short text that can be used to call out experiment records in searches or in displays.

## \*Project accession

Link data to BioProject that describes the research.

## \* Sample accession

Enter a BioSample or GSASample Accession. BioSample accessions have 'SAMN' prefix. A

BioSample describes the biological source material for your sequence library preparation.

## \*Library Construction/Experiment design

Enterthe details about your experimental design and molecular strategies including hybrid

selection and affinity capture reagents; any detail that distinguishes your experiment from

other similar experiments.

### Library

The library descriptor specifies the origin of the material being sequenced and any treatments that the material might have undergone that affect the sequencing result.

### Library name

The submitter's name for this library.

### \*Strategy

Sequencing technique intended for this library.

Strategy	Sequencing strategy used in the experiment	
WGA	Random sequencing of the whole genome following non-PCR	

	amplification		
WGS	Random sequencing of the whole genome		
WXS	Random sequencing of exonic regions selected from the genome		
RNA-Seq	Random sequencing of whole transcriptome		
miRNA-Seq	Random sequencing of small miRNAs		
Tn-Seq	Sequencing from transposon insertion sites		
WCS	Random sequencing of a whole chromosome or other replicon isolated		
	from a genome		
CLONE	Genomic clone based (hierarchical) sequencing		
POOLCLONE	Shotgun of pooled clones (usually BACs and Fosmids)		
AMPLICON	Sequencing of overlapping or distinct PCR or RT-PCR products		
CLONEEND	Clone end (5', 3', or both) sequencing		
FINISHING	Sequencing intended to finish (close) gaps in existing coverage		
ChIP-Seq	Direct sequencing of chromatin immunoprecipitates		
MNase-Seq	Direct sequencing following MNase digestion		
DNase-Hypersensitivity	Sequencing of hypersensitive sites, or segments of open chromatin that		
	are more readily cleaved by DNasel		
Bisulfite-Seq	Sequencing following treatment of DNA with bisulfite to convert		
	cytosine residues to uracil depending on methylation status		
EST	Single pass sequencing of cDNA templates		
FL-cDNA	Full-length sequencing of cDNA templates		
CTS	Concatenated Tag Sequencing		
MRE-Seq	Methylation-Sensitive Restriction Enzyme Sequencing strategy		
MeDIP-Seq	Methylated DNA Immunoprecipitation Sequencing strategy		
MBD-Seq	Direct sequencing of methylated fractions sequencing strategy		
OTHER	Library strategy not listed (please include additional info in the "design		
	description")		

# \*Source

The library source specifies the type of source material that is being sequenced.

Source	Type of genetic source material sequenced		
GENOMIC	Genomic DNA (includes PCR products from genomic DNA)		
TRANSCRIPTOMIC	Transcription products or non-genomic DNA (EST, cDNA, RT-PCR,		
	screened libraries)		
METATRANSCRIPTOMIC	Transcription products from community targets		
METAGENOMIC	Mixed material from metagenome		
SYNTHETIC	Synthetic DNA		
VIRAL RNA	Viral RNA		
OTHER	Other, unspecified, or unknown library source material (please include		
	additional info in the "design description")		

## \*Selection

Whether any method was used to select and/or enrich the material being sequenced.

Selection	Method of selection or enrichment used in the Experiment		
unspecified	Library enrichment, screening, or selection is not specified (please		
	include additional info in the "design description")		
RANDOM	Random selection by shearing or other method		
PCR	Source material was selected by designed primers		
RANDOM PCR	Source material was selected by randomly generated primers		
RT-PCR	Source material was selected by reverse transcription PCR		
HMPR	Hypo-methylated partial restriction digest		
MF	Methyl Filtrated		
CF-S	Cot-filtered single/low-copy genomic DNA		
CF-M	Cot-filtered moderately repetitive genomic DNA		
CF-H	Cot-filtered highly repetitive genomic DNA		
CF-T	Cot-filtered theoretical single-copy genomic DNA		
MDA	Multiple displacement amplification		
MSLL	Methylation Spanning Linking Library		
cDNA	complementary DNA		

ChIP	Chromatin immunoprecipitation		
MNase	Micrococcal Nuclease (MNase) digestion		
DNAse	Deoxyribonuclease (MNase) digestion		
Hybrid Selection	Selection by hybridization in array or solution		
Reduced Representation	Reproducible genomic subsets, often generated by restriction		
	fragment size selection, containing a manageable number of loci to		
	facilitate re-sampling		
Restriction Digest	DNA fractionation using restriction enzymes		
5-methylcytidine antibody	Selection of methylated DNA fragments using an antibody raised		
	against 5-methylcytosine or 5-methylcytidine (m5C)		
MBD2 protein methyl-CpG	Enrichment by methyl-CpG binding domain		
binding domain			
CAGE	Cap-analysis gene expression		
RACE	Rapid Amplification of cDNA Ends		
size fractionation	Physical selection of size appropriate targets		
Padlock probes capture	Circularized oligonucleotide probes		
method			
Poly-A	polyA enriched RNA-seq		
other	Other library enrichment, screening, or selection process		

## \*Layout

Library Layout specifies whether to expect single, Pair-end, or other configuration of reads. In the case of paired reads, information about the relative distance and orientation is specified.

- Fragment
- Paired read

\*Insert size (bp)

Fragment size for Paired reads.

Nominal size (bp)

Size of the insert for Paired reads.

### Nominal standard deviation (bp)

Standard deviation of insert size (typically ~10% of Nominal Size)

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## Run

## **General info**

\* Alias

Submitter assigned name or id for the GSA submission object.

## \* Run data file type Run

The information about supported formats of the submitted sequence data. We recommend that read data is either submitted in Fastq or BAM format. Submitted data files must only contain reads from a single sample.

Format	File suffix	Made available as	Notes
		standard Fastq	
Fastq format	.fastq.gz	Yes	
	.fastq.bz2		
	.fq.gz		
	.fq.bz2		
BAM format	.bam	Yes	
VCF Format	.vcf	No	
SFF Format	.sff	Yes	Spot descriptor is required.

## Data Blocks

## Fastq format

Fastq format is a text-based format for storing both a biological sequence (usually nucleotide sequence) and its corresponding quality scores. Both the sequence letter and

quality score are each encoded with a single ASCII character for brevity.

## \*File Name

We only accept GZIP and BZIP2 compression formats. Especially we don't accept 7-ZIP or TAR compressed files.

- Single reads must be submitted using a single Fastq file and can be submitted the suffix in '\_1', '\_2', etc.
- Paired reads must split and submitted using two Fastq files. The read names must

have a suffix identifying the first and second read from the pair.

Forexample: liver\_Tumor1\_male\_1F.fastq.gz and liver\_Tumor1\_male\_1R.fastq.gz, then followed byread '2F', then '2R', etc.

#### \*MD5 for file

MD5 checksums are a 32-character alphanumeric string. For Mac and Linux system users, the native command line tools "md5sum"(Linux) and "md5"(Mac OX) can be used to generate MD5 checksums. Windows users must need to download a third-party utility.

#### BAM format

The BAM format is an efficient method for storing and sharing data from modern, highly parallel sequencers. While primarily used for storing alignment information, BAMs can (and frequently do) store unaligned reads as well.

### \*Reference Assembly Name

#### \*Assembly Name or Accession

The Reference's assembly name or assembly accession number

## \*Web URL of the Reference Assembly

The URL of the Reference Assembly

#### \*File Name for bam bam

Submitted BAM files must be readable with SAMtools. BAM file names are required to end up with the .bam suffix (e.g. 'a.bam').

#### \*MD5 for bam filebam

MD5 for bam file bam

### \*Local Assembly file

### \*select one reference file you have uploaded/submit new reference file

#### \*Reference file name

The Reference's file name

#### \*MD5 for reference file

MD5 for reference file

## \*File Name for bam

Submitted BAM files must be readable with SAMtools. BAM file names are required to end up with the .bam suffix (e.g. 'a.bam').

### \*MD5 for bam file

MD5 for bam file bam

## VCF format

The Variant Call Format (VCF) specifies the format of a text file used in bioinformatics for storing gene sequence variations. By using the variant call format only the variations need to

be stored along with a reference genome.

### \*Reference Assembly Name

#### \*Assembly Name or Accession

The Reference's assembly name or assembly accession number

## \*Web URL of the Reference Assembly

The URL of the Reference Assembly

### \*File Name forVCF

VCF file names are required to end up with the .vcf suffix (e.g. 'a.vcf').

## \*MD5 for VCF fileVCF

MD5 for VCF file

#### \*Local Assembly file

# \*select one reference file you have uploaded/submit new reference file \*Reference file name

The Reference's file name

\*MD5 for reference file

MD5 for reference file

### \*File Name for VCF

VCF file names are required to end up with the .vcf suffix (e.g. 'a.vcf').

## \*MD5 for VCF file

MD5 for VCF file

## SFF format

Standard flowgram format (SFF) is a binary file format used to encode results of

pyrosequencing from the 454 Life Sciences platform for high-throughput sequencing. SFF

files can be viewed, edited and converted with DNA Baser SFF Workbench (graphic tool), or

converted to FASTQ format with sff2fastq or seq\_crumbs.

#### \*File Name

SFF file names are required to end up with the .sff suffix (e.g. 'a.sff').

## \*MD5 for file

MD5 for SFF file bam

## NOTE:

Transmitting your data files to the GSA FTP site

Address: ftp://submit.big.ac.cn

User: Same as your GSA Username

Password: Same as your GSA Password