## **Genomic Information Database**

Table taken from ISO/IEC JTC 1/SC 29/WG 11 N15092 (Geneva, CH - February 2015)

ID	Sequencing method	Size	File type	Coverage	Origin	Comments	Local path	
Homo Sapiens								
ERP001775	Illumina HiSeq	~2 TB	FastQ	200x	http://www.ebi.ac.uk/ena/data/view/E RP001775	This is the largest dataset. Can be transcoded from gzip to 7zip if needed.	/data/genome/human/illumina/ERP001775	
ERP001960	Illumina HiSeq	~120 GB each genome	BAM	30x	http://www.ebi.ac.uk/ena/data/view/E RP001960	3 genomes selected: SAMEA1573614 SAMEA1573618 SAMEA1573617	/data/genome/human/illumina/ERP001960	
ERP002490	Illumina HiSeq	265 GB	BAM	30-40x	http://www.ebi.ac.uk/ena/data/view/E RP002490	The insert size (distance between the pair of reads) is much larger than usual (about 2K bases instead of 300).	/data/genome/human/illumina/ERP002490	
Low coverage ERR317482 WGS	Illumina HiSeq 2000	6.1 GB	BAM	1.9x	http://www.ebi.ac.uk/ena/data/view/E RR317482 ftp://ftp.1000genomes.ebi.ac.uk/vol1/ ftp/technical/reference/phase2_reference_assembly_sequence/hs37d5.fa.gz	Also used as a low coverage test in the Scramble paper.	/ data/genome/human/illumina/ERR317482W GS	
Low coverage NA21144.ch rom11	Illumina HiSeq 2000	1 GB	BAM	7.5x	ftp://ftp.1000genomes.ebi.ac.uk/vol1/ftp/data/NA21144/alignment/NA2114 4.chrom11.ILLUMINA.bwa.GIH.low_coverage.20130415.bam ftp://ftp.1000genomes.ebi.ac.uk/vol1/ftp/technical/reference/phase2_reference assembly sequence/hs37d5.fa.gz	Used in Scramble paper. Processed through the GATK pipeline, which makes the auxiliary data bulkier. (Can be stripped off easily if desired.)	/ data/genome/human/illumina/NA21144.chro m11	
PacBio_CH M1htert_54x	Pacific Biosciences SMRT	150 GB	FastQ	54x	http://datasets.pacb.com/2014/Human 54x/fast.html	Available as zipped FastQ.	/data/genome/human/pacbio	
IonTorrent	Ion Torrent	1.3 GB each file	BAM		http://www.ebi.ac.uk/ena/data/view/E RX276880 http://www.ebi.ac.uk/ena/data/view/E RX276881 http://www.ebi.ac.uk/ena/data/view/E RX276882	http://www.ebi.ac.uk/ena/data/wareho use/search?query= %22instrument_platform= %22ION_TORRENT %22%20%20AND %20submitted_format=%22BAM %22%22&domain=read	/data/genome/human/IonTorrent	

Author: Jan Voges

RNAseq		16 GB	BAM	http://www.ebi.ac.uk/arrayexpress/file s/E-MTAB-1728/K562_ cytosol_LID8465_TopHat_v2.bam		/data/genome/human/RNASeq		
Bacteria								
ERX593919 (E. Coli)	Oxford Nanopore	60 GB	Gzipped Fast5	http://www.ebi.ac.uk/ena/data/view/E RX593919	To be converted from Fast5.			
ERX593921 (E. Coli)	Oxford Nanopore	46 GB	Gzipped Fast5	http://www.ebi.ac.uk/ena/data/view/E RX593921	To be converted from Fast5.	/data/genome/bacteria/oxfordnano		
DH10B (E.Coli)	Illumina	1.3 GB	BAM	ftp://webdata:webdata@ussd- ftp.illumina.com/Data/SequencingRu ns/DH10B/MiSeq_Ecoli_DH10B_11 0721_PF.bam https://raw.githubusercontent.com/all anroscoche/PathTree/master/data/DH 10B_WithDup_FinalEdit_validated.fa sta	Used in the Deez paper.	/data/genome/bacteria/DH10B		
ERA269036 9799_7#3.ba m (E.Coli)	Illumina	2.3 GB	BAM	ftp://ftp.sra.ebi.ac.uk/vol1/ERA269/E RA269036/bam/ http://www.ncbi.nlm.nih.gov/nuccore/ NC_000913.2? report=fasta&format=text	Used in the Scramble paper.	/data/genome/bacteria/ERA269036		
				Metagenomic				
Human gut	Illumina Genome Analyzer II	10 GB	FastQ	http://www.ebi.ac.uk/ena/data/view/E RA000116	3 samples picked: SAMEA728920 SAMEA728635 SAMEA728854	/data/genome/metagenomic/humangut		
Cancer cell lines								
Mutation/Var iation Calling Benchmark 4 at CGHub		255 GB	BAM	60x https://cghub.ucsc.edu/datasets/bench mark_download.html	TCGA BENCHMARK CELL LINE: HCC1143 NORMAL 60x f0eaa94b-f622-49b9-8eac- e4eac6762598	/data/genome/cancercells/f0eaa94b-f622- 49b9-8eac-e4eac6762598		
Mutation/Var iation Calling		305 GB	BAM	50x https://cghub.ucsc.edu/datasets/bench mark_download.html	TCGA BENCHMARK CELL LINE: HCC1143 TUMOR 50x ad3d4757-f358-40a3-9d92-	/data/genome/cancercells/ad3d4757-f358- 40a3-9d92-742463a95e88		

Last modified: 2015-04-15

Benchmark 4 at CGHub						742463a95e88		
Mutation/Var iation Calling Benchmark 4 at CGHub		130 GB	BAM		https://cghub.ucsc.edu/datasets/bench mark_download.html	UCSC ARTIFICIAL MIXED SAMPLE: 80% HCC1954BL 20% HCC1954 360b4736-6c5e-48df-af58- c1cf51609350	/data/genome/cancercells/360b4736-6c5e- 48df-af58-c1cf51609350	
Plants								
T. Cacao	Illumina	8.2 GB	FastQ	10x	http://www.ncbi.nlm.nih.gov/sra/SRX 288435			

Author: Jan Voges