

Using modern high-throughput sequencing methods, hundreds of thousands of genomic regions can be analyzed in parallel within a very short time and with very high sensitivity.

Next generation sequencing, or high-throughput sequencing, today allows the genetic analysis of the entire genome or transcriptome, a genetic profiling of a person or organism. The possibility of parallel high-throughput sequencing of the entire genome opens up many possibilities in a reasonable time. In addition to whole genome or transcriptome sequencing, methods such as exome sequencing are an alternative to cover all coding regions. Targeted sequencing allows the investigation of disease-associated genes or addresses specific questions. The spectrum of methods is broad and continues to evolve.

- Blood or bone marrow
- Isolated human cells in an appropriate buffer
- FFPE material (Formalin-Fixed Paraffin-Embedded, slices or rolls)
- Liquid Biopsy
- Extracted DNA or RNA of high quality
- Prepared libraries
- Pools ready for sequencing

Sample preparation:

- Isolation of white blood cells from bone marrow and peripheral blood
- Extraction of genomic DNA and RNA from human cells stored in an appropriate buffer
- Extraction of genomic DNA and RNA from FFPE material (slices or rolls)
- Extraction of genomic DNA from Liquid Biopsy

Next generation sequencing:

- Single amplicon sequencing of a defined set of human genes
- Panel Sequencing – Targeted gene panel sequencing for a variety of available panels as well as customized ones
- The enrichment of the target genes follows the library prep by hybridization capture
- WES - Whole exome sequencing (human)
- WGS - Whole genome sequencing
- WTS - Whole transcriptome sequencing (RNA-Seq)
- Methylation Sequencing
- Illumina Sequencing-by-synthesis on MiSeq or NovaSeq 6000 instruments

Data analysis:

- Demultiplexing
- Alignment
- Variant Calling
- Interpretation

Data visualization:

- Quality assessment plots
- Explorative data visualization
- Integrative data visualization

Data transfer:

- Data will be provided as report, excel file or by a link for download

Data security:

Raw sequencing data from the NovaSeq system is directly streamed into a private AWS instance of Amazon Cloud in Frankfurt (AWS, Amazon Web Services) with restricted access. The data is completely anonymized with an arbitrary internal identifier and no personal or clinical data is stored in the cloud. The data security measures comply with the highest standards of the new EU General Data Protection Regulation (GDPR), which has also been verified by external auditors in their reports, including ISO 27001, ISO 27017 and ISO 27018. Furthermore, AWS has also been awarded the C5 attestation of the Federal Office of Information Security. Raw sequencing data from the MiSeq systems is stored locally without external access.

Details:

Sample preparation

- Isolation and enrichment of white blood cells

Enrichment of white blood cells / leukocytes by lysis of the erythrocytes. Subsequently, cells (5 million each) are aliquoted in RLT buffer and can be stored at -80°C.
- DNA extraction from cells

DNA extraction from cells is performed using the Roche MagNA Pure System. The extracted DNA is adjusted to 25ng/μl in 260μl maximum. The storage takes place in tubes with 2D barcode in the Liconic freezing system.
- RNA extraction from cells

RNA extraction from cells is performed using the Roche MagNA Pure System. The quality of the RNA is checked by determination of the concentration and the RIN value. The extracted RNA is stored at -80°C in tubes with 2D barcode in a Liconic freezing system.
- cDNA synthesis

The RNA is then reverse transcribed (RT) using VILO IV transcriptase. RNA of an aliquot/asservat is completely used for RT. The obtained cDNA (50μl) is stored at -20°C in tubes with 2D barcode in a Liconic freezing system.
- Storage of cells

Storage of aliquoted leukocytes at -80°C. One aliquot contains 5 million cells in RLT buffer. The storage takes place in Fluidx tubes identified by a 2D barcode at the bottom of the tube in the Liconic freezing system. The unit of account is the number of tubes/year. The assignment of the sample to the barcode will be announced by MLL when the material is returned.
- Storage of nucleic acid (DNA/RNA/cDNA)

The DNA/RNA/cDNA is stored at the appropriate temperature in tubes with 2D barcodes in a Liconic freezing system. DNA and cDNA are stored at -20°C, RNA at -80°C. The storage takes place in Fluidx tubes identified by a 2D barcode at the bottom of the tube in the Liconic freezing system. The unit of account is the number of tubes/year. The assignment of the sample to the barcode will be announced by MLL when the material is returned.

Library preparation

- Library prep for enrichment (TruSeq DNA Nano)

The library prep for enrichment is performed with a modified Illumina TruSeq DNA Nano using Unique Dual Indices (UDI). The DNA is fragmented to a length of 150bp using the Covaris. The starting material is 150ng DNA. A quality check of the starting DNA is necessary for FFPE or Liquid Biopsy DNA, as it may not have to be fragmented.
- Library prep for enrichment (Illumina DNA Prep)

The library prep for enrichment is performed with the Illumina DNA Prep for Enrichment Kit using Unique Dual Indices (UDI). The DNA is fragmented enzymatically in 180bp fragments in the protocol. 100ng DNA with sufficient fragment length is required as starting material (quality check required for FFPE DNA).
- Hybridization capture based enrichment (gene panel)

The DNA target regions are enriched using the IDT Hybridization Capture Protocol and corresponding xGen lockdown gene panels.
- Library prep for WGS (TruSeq DNA PCR-Free)

The library prep for Whole Genome Sequencing (WGS) is performed with the Illumina TruSeq DNA PCR-Free Kit using Unique Dual Indices (UDI). The DNA is fragmented to a length of 350bp using the Covaris. The starting material is 1µg high-molecular DNA.
- Library prep for WGS with limited material (TruSeq DNA Nano)

The library prep for Whole Genome Sequencing (WGS) is performed with the Illumina TruSeq DNA Nano Kit using Unique Dual Indices (UDI). The DNA is fragmented to a length of 350bp using the Covaris. The starting material is 100ng high-molecular DNA.
- Library prep for WGS/Targeted Seq with fragmented material (xGen cfDNA & FFPE)

The library prep for Whole Genome Sequencing (WGS) is performed with the IDT xGen cfDNA & FFPE Library Prep Kit using Unique Dual Indices (UDI). The starting material is already fragmented DNA from FFPE or Liquid Biopsy, <250ng DNA.
- Library Prep for WTS (TruSeq Stranded Total RNA)

The library prep for Whole Transcriptome Sequencing (WTS) is done with the Illumina TruSeq Stranded Total RNA Kit using Unique Dual Indices (UDI). The starting material is 250ng RNA with a RIN value >5.

- Library prep for RNA-exome sequencing (TruSeq Stranded Total RNA)
The library prep for RNA-exome sequencing is performed with the Illumina TruSeq Stranded Total RNA Kit using Unique Dual Indices (UDI). The starting material is 250ng RNA. FFPE RNA can also be used with this protocol, which usually shows a significantly lower RIN value (RIN <5). The enrichment of the exonic regions is done with the xGen Exome Research Panel v2 and hybridization capture workflow (IDT).
- Library prep for Methylation sequencing (TruSeq Methyl Capture EPIC Library Prep Kit)
The library prep for Methylation sequencing is performed with the Illumina TruSeq Methyl Capture EPIC Library Prep Kit using 24 different indices. The starting material is 500ng human genomic DNA. This protocol is not FFPE compatible.

Sequencing

- Sequencing gene-panel
All targeted NGS analyses will be performed with a mean coverage above 1,000-fold and a sensitivity of 3%. Result reports will be provided as an Excel file providing read depth (mean coverage), DNA annotation, protein annotation, allele frequency [%] of variants, as well as interpretation of the variants in terms of SNP, variant or somatic mutation.
- Sequencing WES (100x)
Sequencing of a WES library with a mean target coverage of 100x on an Illumina NovaSeq 6000 system.
- Sequencing WGS (30x-90x)
Sequencing of a WGS library with a mean target coverage of 30 - 90x on an Illumina NovaSeq 6000 system.
- Sequencing WTS (50 M reads)
Sequencing of a RNA exome library with a target read/fragment count of 50 million on an Illumina NovaSeq 6000 system.
- Sequencing per cluster/read (Mio)
Sequencing of a library/pool based on the targeted readout. The specifications are based on the target read number/cluster passed filter. Sequencing is performed on an Illumina NovaSeq 6000 or MiSeq system. Used adaptor sets have to be compatible with Illumina TruSeq or Nextera adaptors, if not a complete flow cell or lane is requested.
- Sequencing per Gb output
Sequencing of a library/pool based on the targeted Output [Gb]. Sequencing is performed on an Illumina NovaSeq 6000 or MiSeq system. Used adaptor sets have to be compatible with Illumina TruSeq or Nextera adaptors, if not a complete flow cell or lane is requested.

Available Flow Cells/Reagent Kits

- MiSeq instrument
- NovaSeq instrument (SP, S1, S2, S4)
- Lane of NovaSeq S1, S2 or S4 flow cell with 200 or 300 cycles

Data analysis

- Alignment WGS data
Map read data (fastq) to the human reference genome (GRCh37 (hg19), optionally GRCh38) using Isaac aligner, including PCR duplicate marking and indel realignment. Output is a BAM file.
- Alignment RNA Seq data
Map read data (fastq) to the human reference genome (hg19 default, hg38 optionally) using STAR aligner and FPKM and count estimation of reference genes using Cufflinks 2 (transcript estimates with salmon are optionally).
- Tumor/Normal Pipeline for variant calling (SNV/SV/CNV)
BAM (GRCh37, GRCh38) used to detect somatic variants from tumor and matched normal pair using Strelka2 for SNV and small indels, Manta for structural variant calling and GATK4 for CNV. Output is a VCF.
- Tumor only Pipeline for variant calling (SNV/SV/CNV)
BAM (GRCh37, GRCh38) used to detect somatic variants from tumor-only sequencing using Isaac Variant caller for SNV and small indels, Manta for structural variant calling and GATK4 for CNV. Output is a VCF.
- Expression analysis
STAR Aligner BAM is used to perform differential expression analysis of reference genes with edgeR. Control samples or sample grouping (n >= 3) are needed for this analysis.
- Variant calling RNA-Seq: fusion transcripts and SNV
BAM (hg19, hg38) created by STAR aligner. Fusions are called with Arriba, Manta and STAR-Fusion. Fusions called by at least two callers are reported. PE reads required for fusion detection. Additionally Isaac Variant caller is used for SNV and small indel detection.
- Annotation of SNV with public data bases
Annotation of VCF file with Nirvana engine using the following sources: VEP, ClinVar, COSMIC, dbSNP, gnomAD, DGV. Outputs VCF, including alignment and variant calling metrics.
- Manual annotation of SNV by MLL pipeline
Manual annotation of variants using MLL routine diagnostics workflow with variant classification for certain gene panels. Output is a report/excel file.

Data visualization

- Quality assessment plots The visualization of the obtained data set can help to assess the quality of the data, to identify outliers, potential sample swaps, and the performance of data pre-processing and normalization techniques.
- Explorative data visualization Due to the high dimensionality of high-throughput data, unsupervised exploratory analysis techniques are used to obtain a first insight into the dataset.
- Integrative data visualization Genomic data comprises a magnitude of information, and due to its high complexity, there is a need for sophisticated visualization methods and tools to generate new hypothesis and draw conclusions