

Research & Guideline for Transcriptome Sequencing



Guideline for Transcriptome Sequencing

Next-generation sequencing allows the detection and relative quantification of RNA molecules. Transcriptome sequencing is a powerful tool to analyze RNAs in a specific tissue and at a specific time point. It can be used for various purposes.

Transcriptome Workflow

Our transcriptome workflow enables reliable and accurate insights into each step of our process, from RNA extraction to data analysis. Due to permanent and strict quality control as well as validated processing procedures, our clients benefit from high-quality results. Our pipeline is subjected to permanent improvement to offer the best solution possible for all kinds of samples.

Selection of the Appropriate Library Preparation Kit

Transcriptome sequencing protocols are continuously improved at CeGaT. Our goal is to provide our clients with optimal data quality for all applications. Transcriptome sequencing is highly dependent on the quantity and quality of the RNA, and very sensitive to even minute changes in the experimental setup. Considering these, we choose the most suitable protocol for library preparation to achieve the best results possible. Reproducible, comparable and unbiased data are crucial for the comparison within a client's project. Therefore, we always use the same library workflow and have automated processes wherever possible to avoid batch effects.

Our library preparation protocols and wet-lab processes are optimized to generate sequencing libraries using a sequencing read length of 100bp in a paired-end mode (PE100). The sequencing depth is determined by the research objective. Considering the guidelines from the ENCODE (ENCyclopedia of DNA Elements) consortium, we recommend a sequencing depth of 30 M (Million) clusters for mRNA sequencing of human samples. However, this recommendation can differ depending on the transcriptome size of the analyzed organism. Bacteria, for example, require much fewer reads than a hexaploidic plant. To discover low copy number transcripts in samples of particularly low quality and low input, we recommend sequencing at least 50 million clusters.



Step 1 RNA extraction and initial quality control



Step 2

Library preparation including mRNA enrichment or rRNA depletion



Step 3 Next-generation sequencing (NovaSeq™ X Plus & NovaSeq™ 6000)

Step 4

Bioinformatic analysis



Figure 1| Description of the transcriptome workflow. Either the RNA is isolated from primary samples, or directly enters our process. After the RNA passed our internal quality control pipeline, the library preparation with one selected kit for mRNA or total RNA can be started. The resulting libraries are sequenced on Illumina's NovaSeq^{1™} X Plus, NovaSeq^{1™} 6000 and the raw data are processed.

Coding Transcriptome Sequencing versus Whole Transcriptome Sequencing

Most applications for RNA sequencing can be accomplished either by coding transcriptome sequencing (mRNA sequencing) or whole transcriptome (total RNA) sequencing. The starting material for both analyses is single-stranded RNA. We are happy to offer the best method available for the respective research objective.

Coding Transcriptome Sequencing - mRNA Analysis

Sequencing of coding RNA is the most common transcriptome sequencing approach. Its main objective is the quantification of gene expression and differential gene expression analysis, for example the analysis of coding regions (figure 2).

About 1%-5% of the transcriptome corresponds to mRNA (messenger RNA). This type of RNA is mostly poly-adenylated. By targeting all poly-adenylated transcripts, most protein-coding RNA (mRNA) can be enriched. Poly-A enrichment increases the mRNA content and decreases other unwanted RNA molecules present in a total RNA sample. Depending on the protocol, this procedure can result in a coverage of over 85% of bases aligning to coding sequences and UTRs (untranslated regions). This results in an increased sequencing depth on mRNA, including low-level expressed mRNA transcripts, and enables researchers to identify rare transcripts.

Whole Transcriptome Sequencing – Total RNA Analysis

Whole transcriptome sequencing includes the analysis of mRNA, splicing patterns, regulatory regions as well as many forms of non-coding RNAs such as IncRNA and miRNA. It allows a comprehensive view of all transcripts. Ribosomal RNA (rRNA), is usually not the research focus. For better analysis results, rRNA depletion is recommended. Since the efficiency of rRNA depletion has a major impact on the analysis results, CeGaT offers library preparation methods with the most effective rRNA depletion.

Strand Specificity

A strand-specific protocol allows unambiguous assignment of reads to genes that overlap but are located on opposite strands (figure 3). It is impossible to determine the origin of the DNA strand for a certain RNA transcript, if a non-strand-specific protocol has been used.

For coding transcriptome sequencing and whole transcriptome sequencing we offer strand-specific protocols. Identification of the DNA strand, from which the analyzed RNA transcript originated, allows more accurate transcript annotation and the detection of antisense transcript expression.

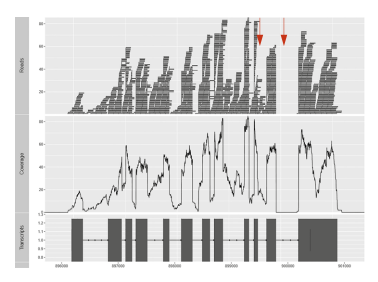


Figure 2 | Representative mapping of a representative gene. The transcripts are shown in the bottom graph and the corresponding coverage is shown in the middle. The read stacks are depicted in the uppermost graph. The coverage is decreased or absent in the intergenic regions (arrows).

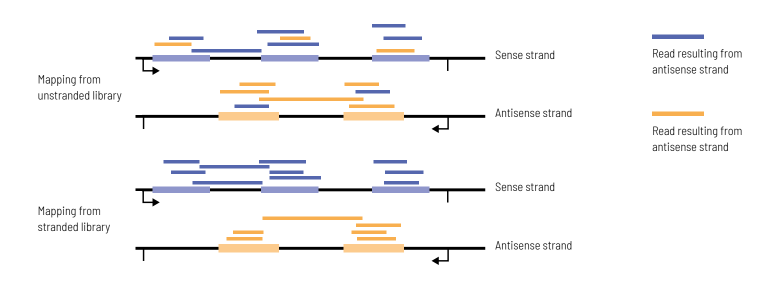


Figure 3 | Comparison of mapping results of sequences obtained from stranded and non-strand-specific libraries. The stranded library enables the assignment of reads to sense and antisense strand.

Transcriptome Sequencing Products

Coding Transcriptome Sequencing (CTS Classic)

CeGaT offers a highly reproducible protocols for a wide range of species for coding transcriptome sequencing. Using our specific mRNA library preparation kit, we offer a consistent workflow with a minimal input of total RNA as starting material (figure 4a). Since mRNA is targeted via its poly-A tail, it is suggested to use good quality RNA with a RIN (RNA Integrity Number) > 8 to ensure unbiased sequencing data. We offer mRNA sequencing for eukaryotes with a recommended output of 30 million clusters per sample using PE100 sequencing.

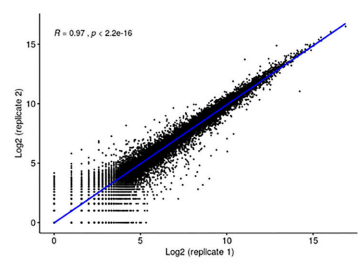
Whole Transcriptome Sequencing (WTS Classic, WTS Deep, TS Flex)

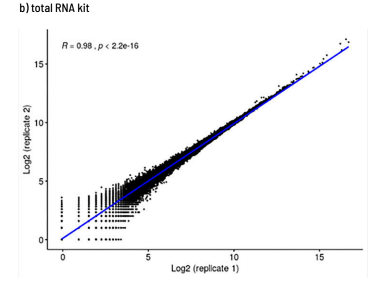
Our total RNA sequencing service provides a reliable and reproducible workflow for analyzing mRNA, as well as many types of non-coding.

RNA and with the option to deplete ribosomal RNA (figure 4b). Even challenging samples with, e.g., moderate or slightly fragmented RNA, RNA of various quality can be used to produce a usable sequencing library. We offer whole transcriptome sequencing for e.g., human, mouse or rat samples. Furthermore, we established workflows to process total RNA samples isolated from human blood, tumors, plants and bacteria. Depending on the sample, RIN values lower than 8 can be accepted. We recommend a sequencing output of 30 million clusters or 50 million clusters, but customization is possible. For more information please get in touch with us.

A considerable part of RNA samples is rather limited and sometimes moderately to highly degraded. We offer adequate solutions of library preparation even for very small amounts of RNA. In case the RNA derived from FFPE tissue or a small number of cells, low input total RNA sequencing is a sensitive solution to accurately detect coding and non-coding RNA transcripts. Our protocol tolerates a DV200 of >30%. Using the low input total RNA kit, CeGaT achieves reproducible results with only 1 ng total RNA of mammalian samples (figure 4c).

a) mRNA kit





c) low input total RNA kit

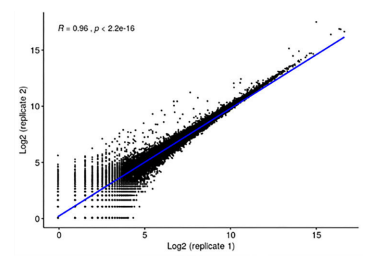


Figure 4 | High correlation of gene expression levels between technical replicates. a) mRNA kit, b) total RNA kit, and c) low input total RNA kit. Normalized counts were used to analyze the reproducibility of gene expression data from UHRR replicate samples.

In-House Data Analysis Services

Always Included:

- ✗ standard data quality control
- $\boldsymbol{\chi}$ detailed project report
- \varkappa delivery of raw data (untrimmed or adapter trimmed FASTQ files)
- $\boldsymbol{\mathcal{X}}$ provision of data via a secure server

Optional:

- * alignment of trimmed sequences for a wide range of eukaryotes, prokaryotes, and viruses (depending on a published reference transcriptome) using STAR (BAM files)
- *x* determination of raw counts per transcript (TSV files)
- × normalization of raw counts using DESeq2 (TSV files; volcano plot, MA plot, PCA plot)
- X statistical comparison of grouped samples to identify differentially expressed genes between different conditions (at least three samples per group; TSV files, heat map)
- X gene ontology (GO) annotation of differentially expressed genes (TSV files)
- X GO term enrichment analysis (TSV files)

By analyzing the normalized read count, differential gene expression analysis can be used to detect quantitative changes in gene expression between experimental groups (figure 5).

Data Security

We operate according to the German Genetic Diagnostics Act. All our data are stored on our servers in-house. We offer end-to-end secured data transfer to our customers and can also provide other individual solutions.

CeGaT guarantees that the data generated within a research project remain the exclusive property of the customer. Samples, sequencing data and analysis results are only used for the purposes specified in the project agreement. We carry out all project steps in-house to ensure no third parties have access to any data. Neither data nor sample material will be sold to others.

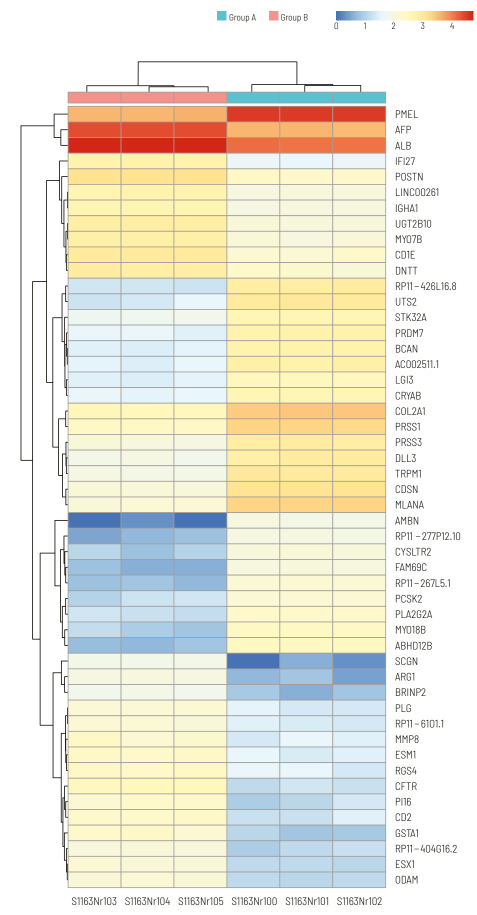
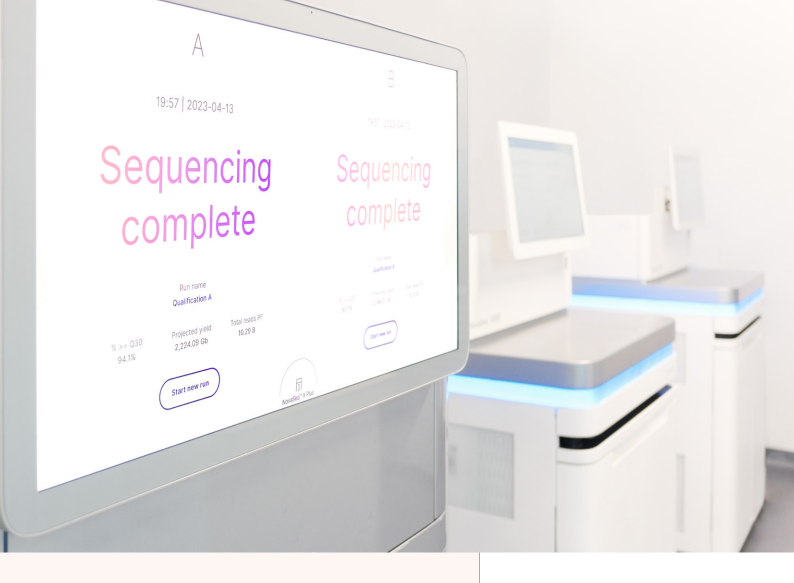


Figure 5 | Gene expression heatmap. This heatmap illustrates the gene expression and group comparison of the top 50 differentially expressed genes. Highly upregulated genes are depicted in red, downregulated genes in blue.



About Us

CeGaT was founded in 2009 in Tübingen, Germany. Our scientists are specialized in next-generation sequencing (NGS) for genetic diagnostics, and we also provide a variety of sequencing services for research purposes and pharma solutions. Our sequencing service portfolio is complemented by analyses suited for microbiome, immunology, and translational oncology studies.

Our dedicated project management team of scientists and bio-informaticians works closely with you to develop the best strategy to realize your project. Depending on its scope, we select the most suitable library preparation and conditions on our sequencing platforms.

We would be pleased to provide you with our excellent service. Contact us today to start planning your next project.



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