

Qlucore Omics Explorer NGS Module feature overview

INTRODUCTION

Qlucore Omics Explorer (QOE) supports the user with fast, simple and visual analysis of measured. The NGS Module is an add-on module that will enable additional functionality related to data generated with NGS technologies and will make it possible to interactively and dynamically analyze and explore NGS data both from DNA and RNA.

All functionality will be provided as integrated parts of Qlucore Omics Explorer and work with the well-known functionality.

MODULE OVERVIEW

The main components of the NGS module are:

- An interactive and fast genome browser for many samples and many tracks per sample
- A genome filter control component
- A project manager for project set-up and definition of which files to include
- A built-in variant caller for short indels and variants
- A Gene Fusions Workbench

INTERACTIVITY

The NGS module is unique in providing true interactive and dynamic analysis of NGS data. The Genome Browser content is dynamically updated when filters and filter cut-off are changed, for instance using sliders and checkboxes.

WORKFLOW AND DATA STORAGE

One unique feature of the NGS Module is the two-stage workflow. The first stage is a pre-processing step that prepares data for the fast and interactive analysis in stage two.

The project files and associated data for indexing samples are stored locally on the user's computer to secure fast access and interactivity while files such as large BAM files can be stored anywhere on a network.

RNA-SEQ ANALYSIS

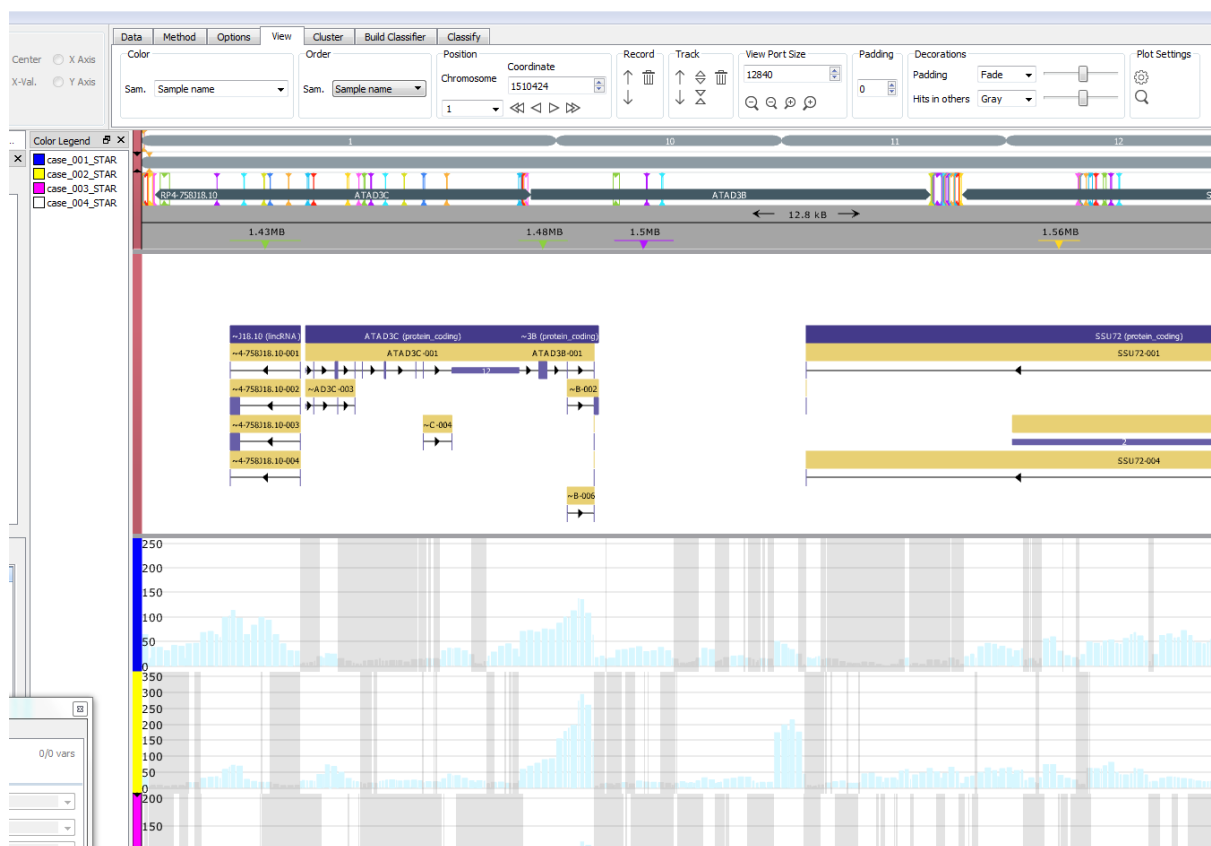
The options available for RNA-seq analysis are impressive. Utilizing the existing functionality in QOE for expression data and combining it with the new NGS functionality enables significantly increased analysis options.

When an RNA-seq project is initiated, both quantitative normalized expression levels (select from FPKM, TPM and TMM) will be calculated at the same time as data is prepared for the genome browser. The program is built to seamlessly handle information both from a quantitative and genomic view. As an example, the user can define discriminating genes with a t-test, visualize them in a heatmap and study the genes in the genome browser. If a sample group is selected or de-selected all plots will be updated and the analysis can directly continue.

GENOME BROWSER

The Genome Browser enables visualization in a variety of flexible ways. Results after filtering, zooming and scrolling are fast and interactive.

A first look at the browser is shown in screenshot below.



Visualization options and selection possibilities are extensive with the fast and interactive genome browser. Examples of options include:

- Display one or several samples

- Add and display any number of annotation tracks
- Detailed track with summary of reads
- Navigation tracks for improved orientation on three levels (whole genome, chromosome and gene)
- Tracks for all accepted data types showing, for example, deletions and insertions
- Tracks with coding regions, genes, introns, exons etc. extracted from GTF file
- Zoom
- Padding to control how much of filtered bases are shown to make sure that edge effects are not affecting the analysis
- Detailed info on selected features in annotations

DATA AND FILE TYPES

A reference genome is mandatory. All organisms are supported, provided a reference genome expressed with the .fasta file format is available.

The following file formats can be imported and used.

- .bam files of version 1.3 or higher. Version 1.5 or higher is¹ recommended
- .vcf files according to VCF version 4.2
- .bed files²
- .gtf files according to specification at <http://www.ensembl.org/info/website/upload/gff.html> ³
- .txt file for cytobands. Format according to UCSC
- .txt or .tsv files for fusion genes. Gene fusion output files from Star Fusion and Fusion Catcher can currently be analysed:
 - Star Fusion: star-fusion.fusion_predictions.abridged.tsv
 - Fusion Catcher: final-list_candidate-fusion-genes.txt

Vcf is a flexible format. A subset of the full VCF file format is supported and the following features are available:

- Handling of small substitutions, insertions and deletions
- Full header parsing
- Support for vcf:s with multiple samples

¹ There is no well-defined standard for a FASTA file. In QOE a FASTA file should have the following structure. A FASTA file contains a list of genomic sequences. Each sequence starts with a header row in this format:

>SequenceID or >SequenceID [Optional Sequence Description]

The header row must start with '>' and be followed immediately (no whitespace in between) by the Sequence ID. The Sequence ID is terminated by whitespace or end-of-line. An optional sequence description can be supplied on the header row - after Sequence ID and the separating white space. The header row is followed by one or more rows of sequence data. Nucleotide sequences are described by A,C,G,T and N characters. Both upper and lower case characters are allowed. The sequence data is terminated by end-of-file, or by a new header line starting with '>'.

² The four first mandatory fields: chr, start, end and name are supported. All further fields are ignored. Track and browser lines are ignored.

³ QOE requires some non-mandatory fields to import files.

- Support for all mandatory SNV fields as well as INFO and FORMAT
- Partial support for large mutations and structural variants

VARIANT CALLING

Variant calling is done using Samtools (<http://www.htslib.org/>) and small variants (insertions, deletions and substitutions) are called by default.

FILTERS

The filter component is flexible and lets the user combine multiple filters to select the relevant parts of data to analyze. The filter can be applied to calculated values such as read coverage as well as presence and matches with certain names. An example is:

Create a combination of filters on parameters such as SNV:s being present in the vcf file and named “A” and not present in the vcf file and named “B” and combine this with a requirement that all displayed bases shall have a read depth of at least 50.

Filtering on any names in GTF files such as gene names, gene ids or transcript ids is supported.

Filtering SNVs on many criteria including INFO or Format values as well and QUAL and FILTER with relevant operators depending on the type of the values is supported.

THE FUSION GENE WORKBENCH AND CIRCLE PLOT

The Workbench have two main tabs, Fusions as detected in the sample(s) and Databases. The Database tab enable inspection of Database content. The program is shipped with the Mitelman and TumorFusions databases. The Fusion tab provides extensive filtering, viewing and sorting options. Examples are:

- Filter on Breakpoint reads
- Filter on Spanning fragments
- Filter on presence in database

Selected Gene fusions can be visualized in the Circle plot.

EXPORTS

- High quality 2-D graphics exports of the genome browser content in png, jpg, bmp, TIFF or pdf (vector graphics)
- Report list of SNV's in the browser and the corresponding information such as BED and Cytoband. HGVS (<http://varnomen.hgvs.org>) strings are provided as well as reference sequence specifiers missing and protein levels

SAMPLE MANAGEMENT

- Samples can be organized into groups using sample annotations
- Active/visible samples can be managed using sample annotations
- Import of sample annotation files (*.txt, *.csv)

SYSTEM REQUIREMENTS

The list below highlights the most important requirements. For full information see the Qlucore Omics Explorer System Requirement..

- 64bit Operating System (Windows or Max OS X)
- 12GB of RAM memory
- A graphical card with support of at least Open GL 3.3.
- At least 500GB free hard disk space, preferable a SSD disk
- A fast processor, preferable Core i7 with at least 4 cores

RESEARCH PURPOSE ONLY

Qlucore Omics Explorer is only intended for research purposes.

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