Qlucore launches Omics Explorer 3.3

Qlucore, a leader in the development of bioinformatics visualization software, has today launched the latest version of its advanced data analysis software. Qlucore Omics Explorer 3.3 includes a new Next Generation Sequencing (NGS) Module and additional new functionalities.

Qlucore Omics Explorer is a next-generation bioinformatics software program for broad usage, known for its easy to use interface and advanced visualization capabilities. Qlucore Omics Explorer 3.3 allows biologists and researchers who are not data analysis experts to explore NGS and data, find new answers and generate new hypotheses.

The NGS Module provides additional functionality related to data generated with NGS technologies which will make it possible to interactively and dynamically visualize, analyze and explore NGS data.

RNA-seq studies is one of the most used techniques in modern genetics and cancer research, and the NGS Module enables a comprehensive and synchronized view of both expression levels and genomic locations. A unique flexibility is handed to the user who can perform a statistical test in the expression space by simply moving a cut-off slider, observing the result in the Genome browser. One of the main components in the NGS Module is the innovative Genome browser which, in combination with the dynamic filters, ensures that the user is always visualizing the most interesting areas of the genome.

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Additional facts about Qlucore Omics Explorer 3.3

Many users in the NGS space today are prevented from undertaking analysis due to the requirements of servers and high performance computers. Qlucore introduces an innovation with assumption free preprocessing. Work is divided in two phases. The pre-process takes time but is only done once and can be run on any local computer or server running Windows or Mac. Visualization and Analysis are performed as a second phase on a normal computer (PC or Mac) which utilizes the pre-processed information.

The main components of the NGS module are:

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

Workflow and data storage

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

RNA-seq analysis

The options available for RNA-seq analysis stand out. Utilization of the existing functionality in QOE for expression data and combining it with the new NGS functionality enable significantly increased options for analysis.

When a RNA-seq project is initiated, the project manager will (during the pre-process stage) calculate both quantitative normalized expression levels as well as prepare the data 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 heat map and at then study the genes in the genome browser. If a sample group is selected or de-selected all samples in both plots will be updated and analysis can continue without any requirements or restarts.

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.

Variant calling

Variant calling can be done with the inbuilt variant caller which uses Samtools (http://www.htslib.org/) or the user can load a VCF file from another variant caller.

Filters

The filter component is flexible and lets the user combine multiple filters to select the relevant regions 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: To create a combination of filters on parameters such as variants being present in the VCF file and named "A" and not present in the VCF file named "B" and combine this with a requirement that all displayed bases shall have a read coverage of at least 50.

Genome browser

The visualization options and possibilities of the "to do" selections are extensive in the fast and interactive genome browser. Example of options are:

- Display one or several samples
- Add and display any number of annotation tracks
- Navigation tracks for improved orientation in three levels (whole genome, chromosome and gene)
- Tracks for all accepted data types showing for instance deletions and insertions
- Tracks with coding regions, genes, introns, exons etc. extracted from GTF file
- Search
- Zoom
- Padding to control how many filtered bases are to be shown to make sure that edge effects are not affecting the analysis

Detailed information on selected features in the annotations

Exports

High quality 2-D graphics exports of the genome browser content in png, jpg, bmp, TIFF or pdf

(vector graphics).

Report list of variants 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)

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About Qlucore

Qlucore (www.qlucore.com) is a leading provider of a new generation intuitive bioinformatics software.

The Qlucore software combines powerful statistical methods with real time visualization. This shortens analysis time, adds more creativity to the research process, strengthens the path to new findings and

facilitates easier and more fruitful collaboration between biologists and bioinformaticians.

The Qlucore Omics Explorer software makes it possible to interactively explore and analyse multivariate

data sets, from small to very large as well as data generated with NGS technologies. The new NGS Module is an add on to Qlucore Omics Explorer, consisting of an interactive Genome browser and

flexible and interactive filter options. One of the key features is to enable synchronization between the

expression analysis of RNA-seq data and genomic information, making researchers even more

productive.

Qlucore was founded in 2007 by leading researchers at the Departments of Mathematics and Clinical

Genetics at Lund University, Sweden. Today Qlucore has customers in about 25 countries around the world, with sales offices in Europe and North America, and distribution in several countries in Asia.

Many of the leading pharmaceutical companies use Qlucore in their research, as well as hospitals and

universities around the world.

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