# **Main Features**

In this section you will find a summary of the main features of OpenCGA.

# Metadata Catalog and Security

OpenCGA Catalog is one of the most important components. Catalog implements the data models, allow custom annotations, implement permissions, ... An audit system has also been implemented.

## **Catalog Data Models and Annotations**

- Rich data models implemented for studies, files, samples, individuals, families, ...
- Advanced free data model implemented for storing custom annotations such as stats or clinical data from patients. Users can define confidential annotations only visible for authorised users.

#### **Catalog Database**

- Catalog database has been implemented using MongoDB to provide a high-performance and s calable query engine.
- Catalog can use Solr as a secondary index to calculate complex annotations and stats.

#### **Authentication and Permissions**

- OpenCGA comes with a built-in authentication system. Other systems are also supported such as LDAP or Microsoft Azure AD (*under development*). Authentication tokens use JWT standard which facilitate the creation of federated systems.
- Advanced and efficient resource permission system implemented in Catalog. You can define
  different permissions such as VIEW, WRITE or DELETE at study level or at any specific
  document. This allow to share data with other users. More information at Sharing and
  Permissions.

## Alignment Storage

OpenCGA can manage alignment data. BAM files can be indexed and coverage calculated.

#### **Fetching alignments**

- Query indexed BAM files, allowed filters include by region, mapping quality, number of mismatches, properly paired, ...
- GA4GH data model used for alignments
- Google gRPC is used as an alternative to REST (JSON) to improve performance.

#### Coverage

- Coverage can be calculated and stored in a BigWig file.
- Coverage queries at any window size or zoom.

## Variant Storage

OpenCGA provides a framework for implementing *big data* variant storage engines which support: realtime queries, interactive complex data aggregations, full-text search, variant analysis, ... The framework takes care of several common operations such as variant normalisation, sample genotype aggregation, variant stats calculation, variant annotation, secondary indexing or in-memory cache. Two different engines are implemented using NoSQL databases: MongoDB and HBase. A secondary index using Solr is nicely integrated with the two implementations. By implementing variant storage engines with NoSQL databases we ensure a fast response time and high concurrent queries.

#### **Data Management**

- Advanced variant normalisation implemented supporting multi-allelic split or left-alignment of INDELs among others.
- High quality sample genotype aggregation supporting multi-allelic variants, overlapping SNV-INDEL or structural variants. HBase storage engine can aggregate tens of thousands of samples efficiently. Current design and implementation should scale to hundreds of thousands of samples.
- **Dynamic** variant storage, you can add or remove samples dynamically from the variant storage efficiently

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- Rich and efficient variant data model implemented. Variant data model support different studies, file information, samples information and a rich variant annotation. Sample genotypes are efficiently stored to scale to hundreds of thousands of genotypes, this allows to optimise analysis by minimising the disk usage and memory consumption.
- Structural variants are fully supported incliuding SNV, INDEL, insertion, deletions, CNV, ...
- Multi-cohort variant stats supported. Users can define different cohorts (group of samples) and
  precompute and index their variant stats, this allows a real-time queries or aggregations. A
  default cohort called *all* is managed automatically.
- CellBase high-performance variant annotation tool is integrated providing rich variant annotations which are stored and indexed, this allows a real-time queries or aggregations. Variant annotation data is returned with the variants since it is part of the data model. Multiple variant annotation can be stored and fetched.
- Custom variant scores from external analysis tools such as GWAS association can be loaded, indexed and queried by.
- Export variant data in different formats such as VCF or Parquet. You can filter which variants and samples are exported.

#### **Query Engine**

- OpenCGA implements a very sophisticated query engine supporting the combination of more than 25 filters: region, genes, type, file attributes, sample genotypes, consequence types, population frequencies, biotype, conservation scores, variant and gene clinical traits, mode of inheritance, disease panels, ... Full-text search is also implemented.
- Other query options supported such as *include, exclude, limit, skip, count,...*Some basic analysis implemented such as compound heterozygous, *de novo* variants, sex
- imputation, unique variant saturation, ...
  Variant query engine supports filtering by sample clinical data thanks to the integration with *Cat alog.*
- MongoDB or HBase are fully integrated with Solr secondary indexes to provide a real-time query engine for all queries and use cases.

#### **Aggregation and Stats**

- Solr integration allows the execution of complex aggregations (faceted search) interactively. Nested and range aggregations are supported. For instance, you can aggregate variants by chromosome and type over 46 million variants in just 2 seconds: http://bioinfo.hpc.cam.ac.uk /hgva/webservices/rest/v1/analysis/variant/stats?timeout=60000&study=reference\_grch37% 3AUK10K&fields=chromosome%3E%3Etype
- Variant query filters for filtering variants and aggregation analysis can be combined to calculate the aggregation of any variant query result.
- Aggregation stats such as average, median, percentile, min, max, ... are also supported

## **Big Data Analysis**

- Variants can be exported to **parquet** file which is an efficient columnar file format. This parquet file can be used by Hive or Spark big data technologies.
- Some complex analysis such as IBS are implemented using a custom Spark library to extend the number of uses cases supported. Note that this analysis can take some time and Spark is not a highly concurrent technology, therefore this analysis are queued by OpenCGA.
- Variant data model store genotypes efficiently ensuring we can execute analysis with tens of thousands of samples.

#### Performance and scalability

- HBase storage engine have been implemented to provide **real-time queries** and **interactive aggregations** (*faceted*) even with tens of thousands of whole genomes.
- Google gRPC is used as an alternative to REST (JSON) to improve performance.
- Some benchmarks with more than 11,000 whole genomes accounting for 25TB show that we can load more than 2,000 files a day and execute most queries in less than 1-2 seconds in a small Hadoop cluster of 20 nodes.
- You can go to HGVA to test OpenCGA query engine performance. HGVA uses OpenCGA and IVA and load about 700 million unique variants from different human studies.

## **Clinical Analysis**

OpenCGA aims to provide a full solution for Clinical Genomics analysis, this covers patient clinical data, interpretation algorithms and a pathogenic variant database.

## **Clinical Data**

- Catalog can store and index any clinical data model for samples, individuals or families. Models are defined by users.
- User can configure the permission and visibility of clinical data using Catalog permissions.

#### **Clinical Interpretation Analysis**

- Open a patient case study by creating a *clinical analysis*, this contains all the patient and family data from Catalog at that moment, the phenotype to be analysed or the files among other information. A rich interpretation data model has also been modelled – combining GEL and other data models – to capture all the relevant information from the interpretation.
- Complete disease panel management implemented: create, update and delete disease panels. You can also import them automatically from PanelApp (GEL). Updated panels are versioned t o keep track of existing interpreted analysis.
- Several rare disease interpretation analysis implemented such as TEAM or Tiering which is based on GEL RD Tiering tool (Cancer interpretation analysis coming soon). You can use one or more disease panels in the interpretation analysis.
- You can save more than one interpretation analysis result in the *clinical analysis* to create one or more clinical reports.
- Together with a tier classification a semi-automatic ACMG classification has been also implemented.

## **Pathogenic Variant Database**

- Interpreted variants and their variant annotation can be indexed in a high-performance patho genic variant database. Clinical data from *catalog*, the *clinical analysis* and *interpretation* are also indexed together with interpreted variants.
- Real-time queries and complex aggregations have been implemented.

# **RESTful Web Services**

OpenCGA implements more than 150 RESTful web services to allow users to manipulate and query Catalog metadata and data such as *alignment*, *variants* and *pathogenic variants*. REST web services are documented using Swagger, you can see OpenCGA Swagger documentation at http://bioinfo.hpc.cam.ac. uk/hgva/webservices/. To facilitate the usage all of these web services we have implemented different client libraries and a command line (see below in Usability).

REST web services can be grouped in different categories: *Catalog, Alignment, Variant, Clinical* and *Admi n.* 

## Catalog

- Catalog data manipulation, you can create, update, delete change permission of data.
- Advanced search web services to query any resource (file, samples, ...)

#### Alignment

- You can index BAM files to query reads and calculate coverage in BigWig format
- Query endpoint to fetch alignments in GA4GH format from several files. Filters implemented include: region, mapping quality, number of mismatches, number of hits, properly paired, ...

#### Variant

- Query variant endpoint allows to query variants by any variant filter. Full control of which fields are returned
- Aggregation stats implemented.
- Others: fetch old variant annotation, variant study metadata, ...

## Clinical

• Several web services to create clinical analysis, execute interpretations or query pathogenic variant database.

## Admin

Administrative web services, only OpenCGA root user can execute them

## Usability

## **REST Clients**

• Four REST clients have been implemented in different programming language: Java, Pythong, R and JavaScript.

## **Command-line Interface (CLI)**

• A fully functional command-line has been implemented

## Visualisation

# **OpenCGA web catalog**

• Web-based application to query and aggregate metadata from catalog

## IVA

- Web-based application for Intercative Variant Analysis
  Highly customisable
  Plugin oriented

#### **Genome Browser**

Genome browser for NGS