Using the Command Line

Installation

As described in the documentation, HGVA backend is powered by the OpenCGA project. The CLI is distributed with the rest of the OpenCGA code. The OpenCGA code can be cloned in your machine by executing in your terminal. Checkout the latest code (release-1.1.0 branch):

```
$ git clone https://github.com/opencb/opencga.git
$ git checkout v1.3.6
```

Alternatively, you can download tar.gz files with the code for the latest tags/releases of OpenCGA from:

https://github.com/opencb/opencga/releases

Once you have downloaded the code, follow the instructions at the *How to Build* section of the OpenCGA repository:

https://github.com/opencb/opencga

The CLI interface is accessible through the opencga.sh script:

```
cd opencga
cd build
cd bin
opencga/build/bin$ ./opencga.sh
Program:
            OpenCGA (OpenCB)
            1.1.0
Version:
Git commit: f2dace56fcdf491efee8ebb0cb43f981e31c320e
Description: Big Data platform for processing and analysing NGS data
            opencga.sh [-h|--help] [--version] <command> [options]
Usage:
Catalog commands:
        users User commands
     projects Project commands
       studies Study commands
        files File commands
         jobs Jobs commands
   individuals Individual commands
     families Family commands
      samples Samples commands
     variables Variable set commands
      cohorts Cohorts commands
Analysis commands:
   alignments Implement several tools for the genomic alignment analysis
       variant Variant commands
```

The CLI provides *commands*, *subcommands* and *parameters* to access its functionality. *Commands* of most interest for HGVA users are *projects*, *studies*, *cohorts* and *samples*. Please, find below a list of commands which can be of most interest for HGVA user. Further documentation on the OpenCGA CLI can be found at the Command Lines section of the OpenCGA documentation.

Configuration

As previously said, the CLI makes intensive use of the RESTful API. Thus, the only configuration detail needed for the CLI to work is a URL where the Web Services API is hosted. The configuration file *client-configuration.yml* is used for this purpose. You shall find a template of this file at the *build/conf* directory:

Table of Contents:

- Installation
- Configuration
- Examples
 - Getting information about variants
 - Getting information about projects
 - Getting information about studies
 - Getting information about samples
 - Getting information about cohorts

Edit this file with any text editor and set the rest host attribute to "http://bioinfo.hpc.cam.ac.uk/hgva":

```
Configuration file client-configuration.yml

---
## number of seconds that session remain open
sessionDuration: 12000

## REST client configuration options
rest:
   host: "http://bioinfo.hpc.cam.ac.uk/hgva"
   batchQuerySize: 200
   timeout: 30000
   defaultLimit: 2000

## gRPC configuration options
grpc:
```

Examples

host: "localhost:9091"

Getting information about variants

You can query variants by using the *variant* command and *query* subcommand. An extensive list of filtering parameters allow great flexibility on the queries. Please, check inline help provided by *opencga*.sh for further details. For example, get TTN variants from the Genome of the Netherlands study, which is framed within the *reference_grch37* project. We will restrict studies data to those corresponding to GONL. Finally, we will also limit the number of returned results to 3:

```
./opencga.sh variant query --gene TTN --study GONL --limit 3 --of json --output-study GONL
```

Getting information about projects

You can use the command *projects* to query projects data.

For getting all metadata from a particular project you can use the *info* subcommand. For example, getting all metadata for the *cancer_grch37* project:

```
./opencga.sh projects info --project cancer_grch37
```

For getting all metadata from all studies associated to a particular project yo ucan use the *studies* subcommand. For example, getting all studies and their metadata for the *cancer_grch37* project:

```
./opencga.sh projects studies --project cancer_grch37
```

Getting information about studies

You can use the command studies to query studies data.

For getting all available studies and their metadata you can use the *search* subcommand. For example, getting all metadata for all available studies (please note, of special interest will be here the field *alias* which contains the study identifier to be used as an input whenever a study must be passed as a parameter):

```
./opencga.sh studies search
```

For getting summary data from a particular study you can use the *summary* subcommand. For example, getting summary data for study *1kG_phase3* which is framed within project *reference_grch37*:

```
./opencga.sh studies summary --study reference_grch37:1kG_phase3
```

For getting all available metadata for a particular study you can use the *info* command. For example, getting all metadata for study *GONL* which is framed within the project *reference_grch37*:

```
./opencga.sh studies info --study GONL
```

For getting all samples metadata for a given study you can use the *samples* subcommand. For example, getting all samples metadata for study 1kG_phase3 which is framed within project reference_grch37. Please, note that not all studies contain samples data, e.g. GONL, ExAC, among others, only provide variant lists and aggregated frequencies, i.e. no sample genotypes.

```
./opencga.sh studies samples --study reference_grch37:1kG_phase3
```

Getting information about samples

You can use the command samples to query samples data.

For getting all metadata for a particular sample you can use the *info* subcommand. For example, get all metadata for sample *HG00096* of the *1kG_phase3* study which is framed within the *reference_grch37* pro ject:

```
./opencga.sh samples info --sample HG00096 --study reference_grch37: 1\mbox{kG\_phase3}
```

Getting information about cohorts

You can use the cohorts command to query cohorts data.

For getting all samples metadata in a given cohort you can use the *samples* subcommand. For example, get all samples metadata for cohort *GBR* from study *1kG_phase3* which is framed within project *reference_grch37*:

```
./opencga.sh cohorts samples --study reference_grch37:1kG_phase3 --cohort GBR
```