

Datasets and Studies

Data organisation

OpenCGA uses a two-level structure to organise datasets, these are *Projects* and *Studies* and are used to organise HGVA data and metadata:

- **Projects** is the top-level and can contain one or more *studies*. *Projects* are specific for one species and assembly, all *studies* in a *project* are stored and indexed together in the same database and, therefore, they share the variant annotation.
- **Study**, in turn, represents a particular dataset which can contain *samples metadata* and *cohorts*, and obviously all the genomic variants. For example, the *1000 Genomes Project* is defined as a *study* in OpenCGA and belongs to *Reference GRCh37* project. You can also define *cohorts* in the *studies*, they are just a set of samples defined within a study. For example, populations and super-populations within The 1000 Genomes Project are defined as *cohorts*, so EUR, AMR or GBR are examples of *cohorts*.

You can get more information about data organisation at [OpenCGA Catalog Data Management](#). *Projects* and *Studies* have a unique **alias** to ease their usage from the command-line and REST API, you can find more information about how to query data programmatically at [RESTful Web Services and Clients](#). Please, see next section the full list and organisation of the currently available *Projects* and *Studies* (datasets) in HVGVA.

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Datasets

In this sections you can find all datasets loaded in HGVA and how they are organised in *Projects* and *Studies* (see previous section).

Project name (<i>alias</i>)	Studies		HGVA Version (date)	
	Name	Alias	v1 (Dec. 2016)	v2 (Jan. 2018)
Reference GRCh37 (<i>reference_grch37</i>)	1000 Genomes Project GRCh37	<i>1kG_phase3</i>	Phase 3 2016-05	Phase 3 2016-05
	Exome Sequencing Project (ESP6500)	<i>ESP6500</i>	2016-05	2016-05
	Exome Aggregation Consortium (ExAC)	<i>EXAC</i>	0.3.1 2016-05	0.3.1 2016-05
	Genome of the Netherlands (GoNL)	<i>GONL</i>	Release 5 2016-05	Release 5 2016-05
	UK10K Project	<i>UK10k</i>	2016-05	2016-05
	DiscovEHR	<i>DISCOVEHR</i>	-	
	Genome Aggregation Database (gnomAD Exomes)	<i>GNOMAD_EXOMES</i>	-	
	Genome Aggregation Database (gnomAD Genomes)	<i>GNOMAD_GENOMES</i>	-	
	Spanish Medical Genome Project (MGP)	<i>MGP</i>	2016-12	2016-12
Reference GRCh38 (<i>reference_grch38</i>)	1000 Genomes Project GRCh38	<i>1kG_phase3</i>	Phase 3 2016-10	Phase 3 2016-10
	ESP6500	<i>ESP6500</i>	-	
	UK10K Project (*)	<i>UK10K</i>	-	
	DiscovEHR (*)	<i>DISCOVEHR</i>	-	
	Genome Aggregation Database (gnomAD Exomes) (*)	<i>GNOMAD_EXOMES</i>	-	
	Genome Aggregation Database (gnomAD Genomes) (*)	<i>GNOMAD_GENOMES</i>	-	
Cancer GRCh37 (<i>cancer_grch37</i>)	QIMR Berghofer Melanoma	<i>QIMR_Berghofer_Melanoma</i>	2016-12	2016-12
	Chronic Myeloid Leukemia - Russian Academy of Medical Sciences	<i>RAMS_CML</i>	2016-12	2016-12
Platinum (<i>platinum</i>)	Illumina Platinum	<i>illumina_platinum</i>	2015-08	2015-08

(*) Liftover carried out by Genomics England (GEL)

Variant Anotation

Variant annotation was carried out by the [CellBase](#) project. Please, check CellBase documentation for details on additional data sources: [Data sources and species](#)