# BLUEPRINT proposed data submission schemas Draft v0.2.2-f167d4a4c2d8e82b716693dd35ee9986a24ba4fe

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## Chapter 1

## **Data Submission**

#### 1.1 Overview of Data Submission Process

There are four major steps in the data submission process:

- 1. Submit raw sequence data to the European Genome-phenome Archive
- 2. Prepare the BLUEPRINT submission files according to DCC data format specifications
- 3. Verify conformity of the submission files
- 4. Submit files to the DCC Secure FTP server

All submitted data must be based on Human reference genome assembly GRCh37 and Ensembl gene set version 68

When submitting experimental data, please make sure you've already deposited your raw data to the appropriate public data repositories (eg: sequencing reads to EBI EGA) and then populate in your submission files the data elements <code>raw\_data\_repository</code> and <code>raw\_data\_accession</code> with the correct repository and accession number respectively.

## 1.2 Preparing Clinical Data and Analyzed Contents for their submission

Submitted clinical or experimental data files must be from any one of these categories:

- Clinical data
- Copy Number Variations
- Gene Expression
- Exon Junctions
- DNA \*-lation (Methylation, Hydroxy-Methylation, Formylation, etc...)
- Protein-DNA interactions
- Regulatory regions

BLUEPRINT DCC is hosting both clinical data and analyzed contents. Contents must be sent following the textual tabular formats defined below. Files with those contents must also follow the BLUEPRINT DCC file naming convention.

Each submitter must have a unique signing key, provided by DACO and DCC. Each file in a submitted archive must be accompanied by its SHA1 **uncompressed** content digest file, digitally signed with the submitter's signing key.

```
# Signed digest of uncompressed contents, will be dlat-p--001-20120920--mycode.txt.shal
openssl dgst -shal -sign subKey.pem -out dlat-p--001-20120920--mycode.txt.shal \
dlat-p--001-20120920--mycode.txt

# Signed digest of already compressed contents
bunzip2 -c dlat-p--001-20120920--mycode.txt.bz2 | openssl dgst -shal -sign subKey.pem \
-out dlat-p--001-20120920--mycode.txt.shal

# Verification of uncompressed contents using
# signed digest dlat-p--001-20120920--mycode.txt.shal
openssl dgst -shal -verify subKey.pem.pub -signature dlat-p--001-20120920--mycode.txt.shal \
dlat-p--001-20120920--mycode.txt

# Verification of compressed contents
bunzip2 -c dlat-p--001-20120920--mycode.txt.bz2 | openssl dgst -shal -verify subKey.pem.pub \
-signature dlat-p--001-20120920--mycode.txt.shal
```

The procedure to submit analyzed contents to BLUEPRINT DCC also involves first having the raw data used for the analysis in the European Genome-phenome Archive (EGA), as all the metadata entries from the analyzed contents to be stored in BLUEPRINT DCC must point to the original raw data.

#### 1.2.1 File Naming Conventions

Submitted files, containing either clinical data or analyzed experiment contents, must follow next file naming convention

```
featureType-fileType--institutionCode-dateFileCreated--freeField.txt
```

```
featureType-fileType--institutionCode-dateFileCreated--freeField.txt.sha1
```

The file name components are mapped in the next way:

Components	Description	Key		
	Clinical data	cdata		
	Copy Number Variations	cngv		
	Gene Expression	exp		
featureType	Exon Junctions	jcn		
	DNA *-lation (Methylation, Hydroxy-Methylation,	dlat		
	Formylation, etc)			
	Protein-DNA interactions	pdna		
	Regulatory regions	rreg		
	Metadata file	m		
	Primary data file	р		
	Secondary data file	S		
flaTuna	Gene expression file	g		
fileType	Donor file	donor		
	Specimen file	specimen		
	Sample file	sample		
	Donor's Family file	family		
institutionCode	institutionCode Institution submitting data			
dateFileCreated The date on which the file is created		YYYYMMDD (ISO-8601)		
freeField An alphanumeric field (max length of 16 chara		e.g.: mysample, 0B1845J		
	where submitters can put internal codes, file sequence			
	numbers, etc			

Different file types of the same feature type are interrelated, because the data they are storing is intertwined. Specific relations are defined on the documentation of each feature type and their file types. For instance, information stored in a primary data file is related and depends on the data from its corresponding metadata file, and the same happens to secondary data files and primary data files. Metadata file contents are related to clinical data sample files.

#### 1.2.2 Tabular File Structure

The submitted analyzed contents are kept in tab-delimited text files. General comments may be added to the beginning of the file with a hash ('#') prefixed at beginning of each comment line. The first non-comment line is the header containing the names of the columns. Each column corresponds to a data element defined in DCC Submission Tabular Formats specification (Chapter 2).

There is a subset of comment lines used to attach data labels to the text files. These data labels follow the form '##labelName value [value ...]'. Currently acknowledged data labels are:

- **format**: This label is **required**, and its value defines the BLUEPRINT data formatting schema used on the file.
- depends: Although this label is not always required, it is important to validate the data coherence of
  the whole data set, because it ensures related data is not corrupted. The values of this label are the
  file on the same submission this file is related to (for instance, the name of a metadata file), and the
  SHA1 digest value (in its hexadecimal representation) of that file's contents.

There are several ways to generate the SHA1 digest of a file, like libraries in most of the programming languages and command-line tools:

```
# Getting the SHA1 digest value of uncompressed contents using OpenSSL
openssl dgst -sha1 dlat-p--001-20120920--mycode.txt

SHA1(dlat-p--001-20120920--mycode.txt)= 81ae49a7014d2d0260625d3535fa6e2a4a0bc06f

# Getting the SHA1 digest value of uncompressed contents using sha1sum
sha1sum dlat-p--001-20120920--mycode.txt

81ae49a7014d2d0260625d3535fa6e2a4a0bc06f dlat-p--001-20120920--mycode.txt
```

An example file is shown below (note that parts of the lines are omitted for readability):

```
# This is an example of a primary analysis file for simple somatic mutations.
# File name: dlat-p--001-20120920--mycode.txt
#
# And it has its labels
##format 0.2.2-f167d4a4c2d8e82b716693dd35ee9986a24ba4fe
##depends dlat-m--001-20120920--mycode.txt 03366af5145107cc818f4827e86b61dcf998ff29
            ⇒analyzed_sample_id
                                ⇒d_lated_fragment_id
                                                    ⇒chromosome
                                                                 \rightarrow...
                                                                        ⊣note
analvsis_id
→dlat:001:1234ff33
                                                    ∌1
                                                           \rightarrow...
                                                                  →#FF#
→dlat:001:00019878
                                                    ∃1
                                                           \dashv...
                                                                  →#FF#
                                                    ⇒|21
                                                           ⇒...
⇒dlat:001:a712838
                                                                  →#FF#
⇒dlat:001:abebdZZZZZ
                                                                  →#FF#
```

All the declared columns for each file type must be set. Data columns are labeled as identifier or reference (I), required (R), desirable (D) or optional (O). Data providers (i.e. submitters) must put all the efforts in order to provide values for the idref and required data columns. The exception for this rule are the desirable fields, required fields which can be unknown on the first submissions, but in that case the fields these exceptions are properly documented.

There are several possible reasons why a column value (either desirable or optional) has not been provided. Next reserved codes must be used to describe the reason:

Code	Meaning
#FF#	Data not supplied at this time (for future fill)
#NA#	Not applicable for the context of the surrounding knowledge
#V0#	Data verified to be unknown (void, undef, null)
#DE#	Data derived from a required or idref field

Some data columns described in this submission manual contain values used as identifiers on BLUEPRINT DCC (e.g. analysis\_id, regulatory\_region\_id, ...). As such, these identifiers should uniquely identify the entity they are referring (an analysis, a regulatory region, ...), and the identifier's value should be globally unique within a center's data submission. Also, these identifiers should be consistent along the different data submissions and releases. If you have to generate your own identifiers, there are some general recommendations, like using the same prefix for the identifiers of the same kind.

When you are submitting string values for columns which can contain URLs or multiple values delimited by commas, each separate value string, before being joined, should be URI encoded.

#### 1.3 File Submission Procedure

Files with the contents to be submitted, along with their corresponding signed disgest, must be sent in a single tar archive. Either the tar archive or its embedded contents should be submitted compressed, using gzip, bzip2 or xz formats.

The following steps are involved in submitting your project's data files to DCC:

- 1. Contact xxx and notify the DCC of your intent to submit data.
- 2. The DCC will provide an SFTP or Aspera account for uploading your data to the DCC's secure server xxx.
- 3. Prepare an archive containing the set of data files (along with their signed digests) comprising your submission.
- 4. Generate a signed digest of the archive your are going to upload, which is also going to be uploaded.
- 5. Login into your SFTP or Aspera account and upload both the data archive and its signed digest.
- 6. Notify the DCC of your successful upload, so they can start the internal validation and processing.

To be finished/defined

## Chapter 2

## DCC Submission Tabular Formats

### 2.1 Clinical Data Submission File Specifications

#### Overview

There are three required clinical and tissue annotation submission files, and one optional template files:

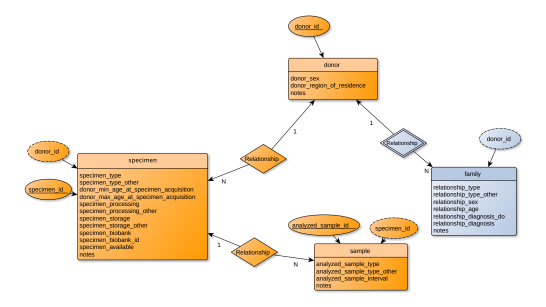


Figure 2.1: Clinical Data Sub-Schema

#### Core Clinical Data Files

- 1. Donor Data File (donor)
  - Mandatory information about the donor's age, gender and vital status.
- 2. Specimen Data File (specimen)
  - **Mandatory** information about a specimen that was obtained from a donor. There may be several specimens per donor that were obtained concurrently or at different times.
- 3. Analyzed Sample Data File (sample)
  - **Mandatory** information about an analyzed sample that was subjected to molecular analysis. There may be several analyzed samples per specimen, for example, when a tumour is used to derive xenografts and cell lines.

All data submissions to the DCC must include the three core clinical data files.

#### **Optional Template Files**

Donor Family History (family)
 Optional details about family history of the donor

#### Coding of donor IDs

The three mandatory data files contain donor, specimen and analyzed sample IDs, respectively. These IDs are to be coded specifically for BLUEPRINT purposes and only the submitting group will keep the key that will permit to link back the data to the individual donors. The key must not be communicated to the data users. It should not be derived from other IDs such as biobank or hospital identifiers. These IDs are to be coded in such a way that they cannot be tracked back to the individual donors, except by the submitting group. IDs are assigned by each submitting group, and must be unique within all the data submitted by that group (i.e. no duplicate IDs allowed). The DCC will prevent collisions between similar IDs submitted by different groups by including the project source column by default in all BioMart queries.

#### 2.1.1 Donor Data File

Donor Data File [donor] (required)

This submission file describes a donor from which one or more specimens were obtained.

Table 2.1.1: Donor Data File

Name	Туре	Need	Description / Values
donor_id	string	I	Unique identifier for the donor; assigned by data provider.
donor_region_of_residence	<pre>string[] (array seps ,)</pre>	R	Country, and optionally state or province code, but not city.  ISO3166-1-alpha-2 or ISO3166-2 codes, eg: "CA" or "CA-ON" (See external CV description A.4)
donor_sex	integer	D	Donor biological sex. "Other" has been removed from the controlled vocabulary due to identifiability concerns.  1 = male 2 = female
notes	string	0	Any additional non-identifying information can be included here.

#### 2.1.2 Donor Family History

Donor Family History [family] (optional)

This file describes the family history of the donor.

Table 2.1.2: Donor Family History

	rable 2.1.2. Bellet 1 aming 1 listery								
Name	Туре	Need	Description / Values						
donor_id	string	I	Unique identifier for the donor; assigned by data provider.						
	Continued on next page								

Table 2.1.2 – concluded from previous page

rable 2.1.2 – concluded from previous page					
Name	Type	Need	Description / Values		
notes str:		0	Any additional non-identifying information can be included here.		
relationship_age integer		R	Age of relative at primary diagnosis (years) Use 90 for ages $>= 90$ years.		
relationship_diagnosis	string	R	Diagnosis (disease or healthy status) e.g. "breast cancer"		
relationship_diagnosis_do	string	R	Disease Ontology code for the relative's diagnosis status		
relationship_sex	integer	R	Biological sex of related individual  1 = male  2 = female		
relationship_type	integer	R	Relationship to the donor  1 = sibling  2 = parent  3 = grandparent  4 = uncle/aunt  5 = cousin  6 = other		
relationship_type_other	string	R	If "other" answered in previous column, specify the re- lationship type here		

#### 2.1.3 Specimen Data File

Specimen Data File [specimen] (required)

This submission file describes a specimen from which one or more samples were derived. Use additional rows for more than one specimen from the same patient. If more than one specimen was extracted during the same procedure, each gets a distinct ID.

Table 2.1.3: Specimen Data File

Name	Туре	Need	Description / Values		
specimen_id	string	I	Unique identifier for the specimen assigned by data provider.		
donor_id	string	I	Unique identifier for the donor; assigned by data provider.		
donor_max_age_at_specimen_acquisition	duration	D	Donor max age when the specimen was acquired. If it is '#DE#', then it is the same value as "donor_min_age_at_specimen_acquisition"		
Continued on next page					

Table 2.1.3 – continued from previous page

Name	Type	Need	Description / Values
donor_min_age_at_specimen_acquisition	duration	R	Donor minimal age when the specimen was acquired, in ISO-8601 duration (basic format)
notes	string	0	Any additional non-identifying information can be included here.
specimen_available	boolean	0	Whether additional tissue is available for followup studies.
specimen_biobank	string	0	If the specimen was obtained from a biobank, provide the biobank name here
specimen_biobank_id	string	0	If the specimen was obtained from a biobank, provide the biobank accession number here.
specimen_processing	string	R	Description of technique used to process specimen  1 = cryopreservation in liquid nitrogen (dead tissue)  2 = cryopreservation in dry ice (dead tissue)  3 = cryopreservation of live cells in liquid nitrogen  4 = cryopreservation, other  5 = formalin fixed, unbuffered  6 = formalin fixed, buffered  7 = formalin fixed & paraffin embedded  8 = fresh  9 = other technique
specimen_processing_other	string	0	If "other" specified for specimen_processing, may indicate technique here.
specimen_storage	string	R	Description of how specimen was stored. For specimens that were extracted freshly or immediately cultured, answer (1) "NA".  1 = frozen, liquid nitrogen 2 = frozen, -70 freezer 3 = frozen, vapor phase 4 = RNA later frozen 5 = paraffin block 6 = cut slide 7 = other
			Continued on next page

Table 2.1.3 – concluded from previous page

Name	Туре	Need	Description / Values
specimen_storage_other	string	0	If "other" specified for specimen_storage, may indicate technique here.
specimen_type	integer	R	Controlled vocabulary description of specimen type.  1 = primary tumour  2 = tumour local recurrence  3 = tumour metastasis to local lymph node  4 = tumour metastasis to distant location  5 = peripheral blood  6 = bone marrow  7 = lymph node  8 = normal control (tissue adjacent to primary)  9 = normal control (blood)  10 = normal control (other)  11 = disease tissue (other)  12 = cord blood
specimen_type_other	string	0	Free text description of site of specimen if "normal control (other)" or "disease tissue (other)" was specified in specimen_type field.

### 2.1.4 Analyzed Sample Data File

Analyzed Sample Data File [sample] (required)

This submission file describes an analyzed sample on which molecular characterization was performed. It includes both control samples and tumour samples.

Table 2.1.4: Analyzed Sample Data File

Name	Туре	Need	Description / Values
analyzed_sample_id	string	I	Unique identifier for the sample assigned by data provider
specimen_id	string	I	Unique identifier for the specimen assigned by data provider.
analyzed_sample_interval	integer	0	Interval from specimen acquisition to sample use in an analytic procedure (e.g. DNA extraction), in days
analyzed_sample_type	integer	R	Controlled vocabulary description of sample type  1 = Normal blood  2 = Leukemic blood  3 = Normal control adjacent to primary  4 = Normal control from non-tumour site  5 = Control from cell line derived from normal tissue  6 = Normal mouse host  7 = Primary tumour  8 = Mouse xenograft derived from tumour  9 = Cell line derived from tumour  10 = Cell line derived from xenograft  11 = Other (specify)
analyzed_sample_type_other	string	0	Free text description of site of sample if "other" was specified in sample_type field
notes	string	0	Any additional non-identifying information can be included here.
purified_cell_type	string	R	Purified cell type for the sample (See external CV description A.3)

## 2.2 Copy Number Germline Variations

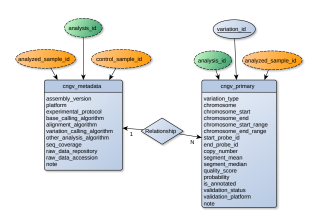


Figure 2.2: Copy Number Germline Variations Sub-Schema

### 2.2.1 Copy Number Germline Variations - Metadata File

Copy Number Germline Variations [cngv] – Metadata File [m]

Table 2.2.1: Copy Number Germline Variations - Metadata File

Name	Туре	Need	Description / Values
analysis_id	string	I	Unique identifier for the analysis performed for a par-
			ticular set of samples
analyzed_sample_id	string	I	Unique identifier for the analyzed sample
alignment_algorithm	compound name;url	R	Name of alignment algorithm and URL to written protocol
assembly_version	integer	R	Version of reference genome assembly (See CV A.8)
base_calling_algorithm	compound name;url	R	Name of base calling algorithm and URL to written protocol
control_sample_id	string	R	Unique identifier for the analyzed control/matched sample
experimental_protocol	compound name;url	0	Name of experimental protocol and URL to written protocol
note	string	0	Optional field to leave notes
other_analysis_algorithm	<pre>compound[]   name;url (array seps ,)</pre>	0	Names of other analysis algorithms. Separate multiple algorithms by commas.
platform	integer	R	Platform or technology used in the detection phase (See CV A.6)
raw_data_accession	compound accession;url	0	Accession and URL for referencing the raw data at the public repository
raw_data_repository	integer	R	Public repository where raw data is submitted (#) (See CV A.9)
seq_coverage	decimal	0	Sequence coverage if analyzed by sequencing plat- forms
variation_calling_algorithm	compound name;url	R	Name of variation calling algorithm and URL to written protocol

## 2.2.2 Copy Number Germline Variations – Primary Analysis File

 $Copy\ Number\ Germline\ Variations\ [cngv]-Primary\ Analysis\ File\ [p]$ 

Table 2.2.2: Copy Number Germline Variations – Primary Analysis File

Tubte	Table 2.2.2. Copy (valide) definiting variations 1 railing / margas 1 te					
Name	Type	Need	Description / Values			
analysis_id	string	I	Unique identifier for the analysis performed for a particular set of samples			
			Continued on next page			

Table 2.2.2 – continued from previous page

Table 2.2.2 – continued from previous page						
Name	Type	Need	Description / Values			
analyzed_sample_id	string	I	Unique identifier for the analyzed sample			
variation_id	string	I	Unique identifier for the variation			
chromosome	integer	R	Name of the chromosome containing the experimentally detected feature (mutation, variation, expression,) (See CV A.7)			
chromosome_end	integer	R	End position of the mutation/variation on the chromosome			
chromosome_end_range	integer	R	Number of bases around chromosome_end that may contain the end position  0 if end position is exactly at chromosome_end; positive integer for +/- number of bases around chromosome_end			
chromosome_start	integer	R	Start position of the mutation/variation on the chromosome			
chromosome_start_range	integer	R	Number of bases around chromosome_start that may contain the start position  0 if start position is exactly at chromosome_start; positive integer for +/- number of bases around chromosome_start			
chromosome_strand	integer	0	Strand where it was detected the mutation/variation on the chromosome -1 = Reverse strand 1 = Forward strand			
copy_number	decimal	0	DNA copy number estimated			
end_probe_id	string	0	Probe id containing the chromosome_end (if array platform was used)			
is_annotated	boolean	0	Indicate if the variation is annotated in the database of Genomic Variations			
note	string	0	Optional field to leave notes			
probability	decimal	0	Probability of the mutation/variation call			
quality_score	decimal	0	Average quality score for the mutation/variation call			
segment_mean	decimal	0	Mean LRR per segment			
segment_median	decimal	0	Median LRR per segment			
			Continued on next page			

Table 2.2.2 – concluded from previous page

Name	Туре	Need	Description / Values
start_probe_id	string	0	Probe id containing the chromosome_start (if array platform was used)
			prationii was useu)
validation_platform	integer	0	Platform or technology used in validation
			(See CV A.6)
validation_status	integer	R	Indicate if the mutation/variation has been validated
			-1 = Not valid
			0 = Not tested
			1 = Validated
variation_type	string	R	Type of variation
			1 = gain
			2 = loss
			3 = copy neutral LOH
			4 = copy neutral
			5 = hemizigous del LOH
			6 = apm LOH

## 2.3 Gene Expression

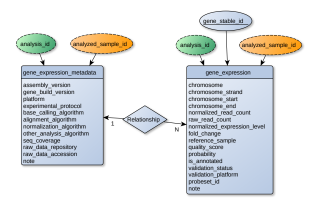


Figure 2.3: Gene Expression Sub-Schema

#### 2.3.1 Expression - Metadata File

Expression [exp] – Metadata File [m]

Table 2.3.1: Expression - Metadata File

Name	Type	Need	Description / Values
analysis_id	string	I	Unique identifier for the analysis performed for a particular set of samples
			Continued on next page

Table 2.3.1 – concluded from previous page

Name	Туре	Need	Description / Values
analyzed_sample_id	string	I	Unique identifier for the analyzed sample
alignment_algorithm	compound name;url	R	Name of alignment algorithm and URL to written protocol
assembly_version	integer	R	Version of reference genome assembly (See CV A.8)
base_calling_algorithm	compound name;url	R	Name of base calling algorithm and URL to written protocol
experimental_protocol	compound name;url	0	Name of experimental protocol and URL to written protocol
note	string	0	Optional field to leave notes
other_analysis_algorithm	<pre>compound[]   name;url (array seps ,)</pre>	0	Names of other analysis algorithms. Separate multiple algorithms by commas.
platform	integer	R	Platform or technology used in the detection phase (See CV A.6)
raw_data_accession	compound accession;url	0	Accession and URL for referencing the raw data at the public repository
raw_data_repository	integer	R	Public repository where raw data is submitted (#) (See CV A.9)
seq_coverage	decimal	0	Sequence coverage if analyzed by sequencing plat- forms

## 2.3.2 Expression - Gene File

 $\mathsf{Expression} \ [\mathsf{exp}] - \mathsf{Gene} \ \mathsf{File} \ [\mathsf{g}]$ 

Table 2.3.2: Expression - Gene File

Name	Туре	Need	Description / Values	
analysis_id	string	I	Unique identifier for the analysis performed for a particular set of samples	
analyzed_sample_id	string	I	Unique identifier for the analyzed sample	
gene_stable_id	string	I	For annotated gene, use Ensembl gene ID. Otherwise, use assemblyBuild_chr_start_end where assemblyBuild is hg18 or hg19.	
Continued on next page				

Table 2.3.2 – continued from previous page

Namo		Need	nued from previous page  Description / Values
Name	Type		
chromosome	integer	R	Name of the chromosome containing the experimentally detected feature (mutation, variation, expression,) (See CV A.7)
chromosome_end	integer	R	End position of the mutation/variation on the chromosome
chromosome_start	integer	R	Start position of the mutation/variation on the chromosome
chromosome_strand	integer	0	Strand where it was detected the mutation/variation on the chromosome -1 = Reverse strand 1 = Forward strand
fold_change	decimal	0	Expressed fold change if differential expression is measured
is_annotated	boolean	R	Indicate if the expressed fragment is annotated in Ensembl
normalized_expression_level	decimal	0	Normalized value of expression level if analyzed by microarray platforms
normalized_read_count	decimal	R	Normalized count of sequencing reads if analyzed by sequencing platforms
note	string	0	Optional field to leave notes
probability	decimal	0	Probability of the mutation/variation call
probeset_id	string	0	ID of the probeset used in microarray if analyzed by microarray platform
quality_score	decimal	0	Average quality score for the mutation/variation call
raw_read_count	integer	R	Raw count of sequencing reads if analyzed by sequencing platforms
reference_sample	string	0	ID of the reference analyzed sample if differential expression is measured
validation_platform	integer	0	Platform or technology used in validation (See CV A.6)
validation_status	integer	R	Indicate if the mutation/variation has been validated -1 = Not valid 0 = Not tested 1 = Validated
			Continued on next page

## Type | Need Description / Values

#### **Exon Junction** 2.4

The following diagram, based on the one from ICGC DCC manual, illustrates how junction\_id should be generated, how junction\_read\_count, exon1\_number\_bases and exon2\_number\_bases are calculated:

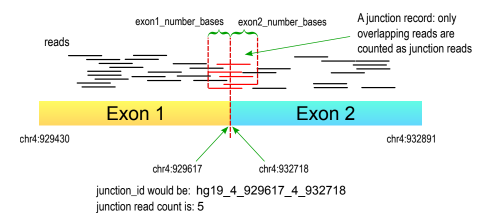


Figure 2.4: Junction Read Count explanation

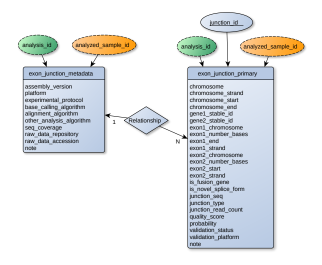


Figure 2.5: Exon Junction Sub-Schema

#### Exon Junction - Metadata File

Exon Junction [jcn] – Metadata File [m]



Table 2.4.1 – concluded from previous page

Name	Type	Need	Description / Values
	3.		<u> </u>
analysis_id	string	Ι	Unique identifier for the analysis performed for a particular set of samples
	-		
analyzed_sample_id	string	I	Unique identifier for the analyzed sample
alignment_algorithm	compound	R	Name of alignment algorithm and URL to written pro-
	name;url		tocol
assembly_version	integer	R	Version of reference genome assembly (See CV A.8)
base_calling_algorithm	compound	R	Name of base calling algorithm and URL to written
	name;url		protocol
experimental_protocol	compound	0	Name of experimental protocol and URL to written
	name;url		protocol
note	string	0	Optional field to leave notes
other_analysis_algorithm	compound[]	0	Names of other analysis algorithms. Separate multi-
3 3	name;url		ple algorithms by commas.
	(array seps ,)		
platform	integer	R	Platform or technology used in the detection phase $(See\ CV\ A.6)$
raw_data_accession	compound	0	Accession and URL for referencing the raw data at the
Taw_uata_accession	accession;url	U	public repository
	accession;urt		
raw_data_repository	integer	R	Public repository where raw data is submitted (#) (See CV A.9)
seq_coverage	decimal	0	Sequence coverage if analyzed by sequencing plat-
			forms

## 2.4.2 Exon Junction - Primary Analysis File

Exon Junction [jcn] – Primary Analysis File [p]

string	I	
	_	Unique identifier for the analysis performed for a particular set of samples
string	I	Unique identifier for the analyzed sample
string	I	For known exons, use exonID1_exonID2 where exonID1 and exonID2 are Ensembl IDs of the 5' and 3' exons, respectively. For novel or putative exons, use assembly—Build_exon1chr_exon1end_exon2chr_exon2start where assemblyBuild is hg18 or hg19; exon1chr and exon2chr are the chromosomes of the 5' and 3' exons, respectively; exon1end is the end position of the 5' exon; exon2start is the start position of the 3' exon.
integer	R	Name of the chromosome containing the experimentally detected feature (mutation, variation, expression,) (See CV A.7)
integer	R	End position of the mutation/variation on the chromosome
integer	R	Start position of the mutation/variation on the chromosome
integer	0	Strand where it was detected the mutation/variation on the chromosome -1 = Reverse strand 1 = Forward strand
integer	R	Name of the chromosome containing the 5' exon (#) (See CV A.7)
integer	R	End position of the 5' exon on the chromosome
integer	R	Number of bases from 5' exon
integer	0	Chromosome strand of the 5' exon -1 = Reverse strand 1 = Forward strand
integer	R	Name of the chromosome containing the 3' exon (#) (See CV A.7)
integer	R	Number of bases from 3' exon
integer	R	Start position of the 3' exon on the chromosome
integer	0	Chromsome strand of the 3' exon -1 = Reverse strand 1 = Forward strand  Continued on next page
	integer	integer R

Table 2.4.2 – concluded from previous page

Name	Туре	Need	Description / Values
gene1_stable_id	string	R	Stable ID of the gene containing the 5' exon at the junction. For annotated gene, use Ensembl gene ID. For putative and novel gene, use assemblyBuild_chr_start_end where assemblyBuild can be hg18 or hg19.
gene2_stable_id	string	0	In the case of a fusion gene, provide the Stable ID of the gene containing the 3' exon at the junction. For annotated genes, use Ensembl gene ID. For putative and novel genes, use assemblyBuild_chr_start_end where assemblyBuild can be hg18 or hg19.
is_fusion_gene	boolean	0	Indicate if the function is the result of a fusion gene
is_novel_splice_form	boolean	0	Indicate if the splice form is novel
junction_read_count	integer	R	Count of sequencing reads that span across exons
junction_seq	string	0	Provide junction sequence if either is_fusion_gene or is_novel_splice_form is true
junction_type	integer	0	Type of junction 1 = Canonical 2 = Non-canonical 3 = U12
note	string	0	Optional field to leave notes
probability	decimal	0	Probability of the mutation/variation call
quality_score	decimal	0	Average quality score for the mutation/variation call
validation_platform	integer	0	Platform or technology used in validation (See CV A.6)
validation_status	integer	R	Indicate if the mutation/variation has been validated -1 = Not valid 0 = Not tested 1 = Validated

### 2.5 Protein-DNA interactions

### 2.5.1 Protein-DNA interaction - Metadata File

 $Protein-DNA \left[pdna\right]-Metadata \ File \left[m\right]$ 

Table 2.5.1: Protein-DNA interaction - Metadata File

Name	Type	Need	Description / Values
analysis_id	string	I	Unique identifier for the analysis performed for a particular set of samples
			Continued on next page

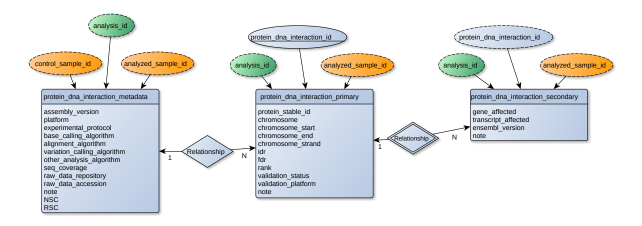


Figure 2.6: Protein-DNA interactions Sub-Schema

Table 2.5.1 – continued from previous page

NI.	_		ed from previous page
Name	Туре	Need	Description / Values
analyzed_sample_id	string	I	Unique identifier for the analyzed sample
NSC	decimal	0	Normalized strand-cross correlation of the analysis
RSC	decimal	0	Relative strand-cross correlation of the analysis
alignment_algorithm	compound name;url	R	Name of alignment algorithm and URL to written protocol
assembly_version	integer	R	Version of reference genome assembly (See CV A.8)
base_calling_algorithm	<b>compound</b> name;url	R	Name of base calling algorithm and URL to written protocol
control_sample_id	string	R	Unique identifier for the analyzed control/matched sample
experimental_protocol	compound name;url	0	Name of experimental protocol and URL to written protocol
note	string	0	Optional field to leave notes
other_analysis_algorithm	<pre>compound[]   name;url (array seps ,)</pre>	0	Names of other analysis algorithms. Separate multiple algorithms by commas.
platform	integer	R	Platform or technology used in the detection phase (See CV A.6)
raw_data_accession	<pre>compound accession;url</pre>	0	Accession and URL for referencing the raw data at the public repository
	l		Continued on next page

Table 2.5.1 – concluded from previous page

Name	Туре	Need	Description / Values
raw_data_repository	integer	R	Public repository where raw data is submitted (#) (See CV A.9)
seq_coverage	decimal	0	Sequence coverage if analyzed by sequencing plat- forms

### 2.5.2 Protein-DNA interaction - Primary Analysis File

Protein-DNA [pdna] – Primary Analysis File [p]

Table 2.5.2: Protein-DNA interaction - Primary Analysis File

Name	Туре	Need	Description / Values	
analysis_id	string	I	Unique identifier for the analysis performed for a particular set of samples	
analyzed_sample_id	string	I	Unique identifier for the analyzed sample	
protein_dna_interaction_id	string	I	Unique identifier for the protein-DNA interaction	
chromosome	integer	R	Name of the chromosome containing the experimentally detected feature (mutation, variation, expression,) (See CV A.7)	
chromosome_end	integer	R	End position of the mutation/variation on the chromosome	
chromosome_start	integer	R	Start position of the mutation/variation on the chromosome	
chromosome_strand	integer	0	Strand where it was detected the mutation/variation the chromosome -1 = Reverse strand 1 = Forward strand	
fdr	decimal	R	False discovery rate	
idr	decimal	R	Irreproducible discovery rate	
note	string	0	Optional field to leave notes	
protein_stable_id	string	R	Stable id of the interacting protein, antibody or protein complex	
rank	<pre>compound[]   rank;value (array seps  )</pre>	0	Kind of used ranking and its value, in the form "rank;value". As it can hold more than one value, they are separated by bars	
			Continued on next page	

Table 2.5.2 – concluded from previous page

Name	Туре	Need	Description / Values
validation_platform	integer	0	Platform or technology used in validation (See CV A.6)
validation_status	integer	R	Indicate if the mutation/variation has been validated -1 = Not valid 0 = Not tested 1 = Validated

### 2.5.3 Protein-DNA interaction - Secondary Analysis File

Protein-DNA [pdna] – Secondary Analysis File [s]

Table 2.5.3: Protein-DNA interaction - Secondary Analysis File

Name	Туре	Need	Description / Values
analysis_id	string	I	Unique identifier for the analysis performed for a particular set of samples
analyzed_sample_id	string	I	Unique identifier for the analyzed sample
ensembl_version	integer	R	Version of Ensembl gene build used for annotation
gene_affected	<pre>string[] (array seps  )</pre>	R	Gene affected. Use Ensembl gene id, separated by   when there is more than one. If no gene is affected, don't put an entry (See external CV description A.1)
note	string	0	Optional field to leave notes
protein_dna_interaction_id	string	I	Unique identifier for the protein-DNA interaction
transcript_affected	<pre>string[] (array seps  )</pre>	0	Transcript on the protein-DNA interaction area. Use Ensembl transcript id. Separate multiple transcripts with vertical bars in the form of transcriptA transcriptB transcriptC (See external CV description A.2)

### 2.6 Regulatory Regions

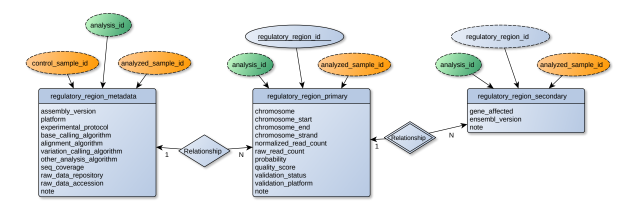


Figure 2.7: Regulatory Regions Sub-Schema

### 2.6.1 Regulatory regions - Metadata File

 $Regulatory\ regions\ [rreg]-Metadata\ File\ [m]$ 

Table 2.6.1: Regulatory regions - Metadata File

Name	Type	Need	Description / Values
			•
analysis_id	string	I	Unique identifier for the analysis performed for a particular set of samples
analyzed_sample_id	string	Ι	Unique identifier for the analyzed sample
alignment_algorithm	compound name;url	R	Name of alignment algorithm and URL to written protocol
assembly_version	integer	R	Version of reference genome assembly (See CV A.8)
base_calling_algorithm	compound name;url	R	Name of base calling algorithm and URL to written protocol
control_sample_id	string	R	Unique identifier for the analyzed control/matched sample
experimental_protocol	<b>compound</b> name;url	0	Name of experimental protocol and URL to written protocol
note	string	0	Optional field to leave notes
other_analysis_algorithm	<pre>compound[]   name;url (array seps ,)</pre>	0	Names of other analysis algorithms. Separate multiple algorithms by commas.
platform	integer	R	Platform or technology used in the detection phase (See CV A.6)
raw_data_accession	<pre>compound accession;url</pre>	0	Accession and URL for referencing the raw data at the public repository
raw_data_repository	integer	R	Public repository where raw data is submitted (#) (See CV A.9)
seq_coverage	decimal	0	Sequence coverage if analyzed by sequencing plat- forms

### 2.6.2 Regulatory regions - Primary Analysis File

Regulatory regions [rreg] – Primary Analysis File [p]

Table 2.6.2: Regulatory regions - Primary Analysis File

Name	Туре	Need	Description / Values
analysis_id	string	I	Unique identifier for the analysis performed for a particular set of samples
			Continued on next page

Table 2.6.2 – concluded from previous page

	Table 2.0.2 – concluded from previous page						
Name	Type	Need	Description / Values				
analyzed_sample_id	string	I	Unique identifier for the analyzed sample				
regulatory_region_id	string	I	Unique identifier for the identified regulatory region				
chromosome	integer	R	Name of the chromosome containing the experimentally detected feature (mutation, variation, expression,) (See CV A.7)				
chromosome_end	integer	R	End position of the mutation/variation on the chromosome				
chromosome_start	integer	R	Start position of the mutation/variation on the chromosome				
chromosome_strand	integer	0	Strand where it was detected the mutation/variation on the chromosome -1 = Reverse strand 1 = Forward strand				
normalized_read_count	decimal	R	Normalized count of sequencing reads if analyzed by sequencing platforms				
note	string	0	Optional field to leave notes				
probability	decimal	0	Probability of the mutation/variation call				
quality_score	decimal	0	Average quality score for the mutation/variation call				
raw_read_count	integer	R	Raw count of sequencing reads if analyzed by sequencing platforms				
validation_platform	integer	0	Platform or technology used in validation (See CV A.6)				
validation_status	integer	R	Indicate if the mutation/variation has been validated -1 = Not valid 0 = Not tested 1 = Validated				

### 2.6.3 Regulatory regions - Secondary Analysis File

Regulatory regions [rreg] – Secondary Analysis File [s]

Table 2.6.3: Regulatory regions - Secondary Analysis File

Name	Type	Need	Description / Values
analysis_id	string	I	Unique identifier for the analysis performed for a particular set of samples
			Continued on next page

Table 2.6.3 – concluded from previous page

Name	Туре	Need	Description / Values
analyzed_sample_id	string	I	Unique identifier for the analyzed sample
ensembl_version	integer	R	Version of Ensembl gene build used for annotation
gene_affected	<pre>string[] (array seps  )</pre>	R	Gene affected. Use Ensembl gene id, separated by when there is more than one. If no gene is affected, don't put an entry (See external CV description A.1)
note	string	0	Optional field to leave notes
regulatory_region_id	string	I	Unique identifier for the identified regulatory region

## 2.7 DNA \*-lation (Methylation, Hydroxy-Methylation, Formylation, etc...)

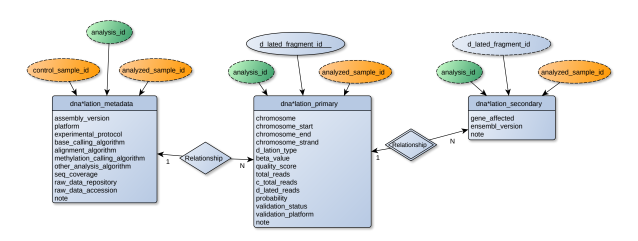


Figure 2.8: DNA Methylation , Hydroxy-Methylation, Formylation, etc... Sub-Schema

#### 2.7.1 DNA \*-lation - Metadata File

DNA \*-lation [dlat] - Metadata File [m]

	Table	2./.	1:	DNA	*-lation -	Metadata	⊦ile
-							

Name	Туре	Need	Description / Values
analysis_id	string	I	Unique identifier for the analysis performed for a particular set of samples
analyzed_sample_id	string	I	Unique identifier for the analyzed sample
			Continued on next page

Table 2.7.1 – concluded from previous page

Name	Type	Need	Description / Values
alignment_algorithm	compound	R	Name of alignment algorithm and URL to written pro-
	name;url		tocol
assembly_version	integer	R	Version of reference genome assembly (See CV A.8)
base_calling_algorithm	compound name;url	R	Name of base calling algorithm and URL to written protocol
control_sample_id	string	R	Unique identifier for the analyzed control/matched sample
experimental_protocol	compound name;url	0	Name of experimental protocol and URL to written protocol
methylation_calling_algorithm	compound name;url	0	Name of variation calling algorithm and URL to written protocol
note	string	0	Optional field to leave notes
other_analysis_algorithm	<pre>compound[]   name;url (array seps ,)</pre>	0	Names of other analysis algorithms. Separate multiple algorithms by commas.
platform	integer	R	Platform or technology used in the detection phase (See CV A.6)
raw_data_accession	<pre>compound accession;url</pre>	0	Accession and URL for referencing the raw data at the public repository
raw_data_repository	integer	R	Public repository where raw data is submitted (#) (See CV A.9)
seq_coverage	decimal	0	Sequence coverage if analyzed by sequencing plat- forms

## 2.7.2 DNA $^*$ -lation - Primary Analysis File

DNA \*-lation [dlat] - Primary Analysis File [p]

Table 2.7.2: DNA  $^*$ -lation - Primary Analysis File

Name	Type	Need	Description / Values
analysis_id	string	I	Unique identifier for the analysis performed for a particular set of samples
analyzed_sample_id	string	I	Unique identifier for the analyzed sample
d_lated_fragment_id	string	I	Unique identifier for the methy- lated fragment, in the form d'*lationType chromosome_chromosomeStart_chromosom
	•		Continued on next page

Table 2.7.2 – continued from previous page

Name			Continued from previous page
	Type	Need	Description / Values
beta_value	decimal	0	DNA *-lation beta value for interrogated site
c_total_reads	decimal	R	Reads which has identified this position as a cytosine
chromosome	integer	R	Name of the chromosome containing the experimentally detected feature (mutation, variation, expression,) (See CV A.7)
chromosome_end	integer	R	End position of the mutation/variation on the chromosome
chromosome_start	integer	R	Start position of the mutation/variation on the chromosome
chromosome_strand	integer	0	Strand where it was detected the mutation/variation on the chromosome -1 = Reverse strand 1 = Forward strand
d_lated_reads	decimal	R	Reads which has identified this position as a DNA *lated cytosine
d_lation_type	string	R	Type of DNA *-lation (Methylation, Hydroxy-Methylation, Formylation, etc)  m = Methylation (cytosine)  hm = Hydroxy-Methylation (cytosine)  hmU = Hydroxy-Methylation (uracil)  f = Formylation (cytosine)  ca = Carboxylation (cytosine)
note	string	0	Optional field to leave notes
probability	decimal	0	Probability of the DNA *-lation call
quality_score	decimal	0	Quality score for the DNA *-lation call
total_reads	decimal	R	Total number of reads over this position/segment, either identifying or not a cytosine, for sequencing platforms. Mean reads depth for other technologies
validation_platform	integer	0	Platform or technology used in validation (See CV A.6)
validation_status	integer	R	Indicate if the mutation/variation has been validated -1 = Not valid 0 = Not tested 1 = Validated
			Continued on next page

# Table 2.7.2 – concluded from previous page Name Type Need Description / Values

## 2.7.3 DNA $^*$ -lation - Secondary Analysis File

DNA \*-lation [dlat] - Secondary Analysis File [s]

Table 2.7.3: DNA \*-lation - Secondary Analysis File

Name	Туре	Need	Description / Values	
analysis_id	string	I	Unique identifier for the analysis performed for a particular set of samples	
analyzed_sample_id	string	Ι	Unique identifier for the analyzed sample	
d_lated_fragment_id	string	I	Unique identifier for the methy- lated fragment, in the form d'*lationType chromosome_chromosomeStart_chromoso	meEnd
ensembl_version	integer	R	Version of Ensembl gene build used for annotation	
gene_affected	<pre>string[] (array seps  )</pre>	R	Gene affected. Use Ensembl gene id, separated by   when there is more than one. If no gene is affected, don't put an entry (See external CV description A.1)	
note	string	0	Optional field to leave notes	

## Appendix A

## Controlled Vocabularies

#### A.1 Ensembl Genes

Valid Ensembl Genes identifiers

(See it at http://jul2012.archive.ensembl.org/Homo\_sapiens/Info/Index)

### A.2 Ensembl Transcripts

Valid Ensembl Transcript identifiers

(See it at http://jul2012.archive.ensembl.org/Homo\_sapiens/Info/Index)

### A.3 Cell Ontology

The Cell Ontology is designed as a structured controlled vocabulary for cell types (See it at http://cellontology.org/)

#### A.4 ISO 3166-1 and ISO 3166-2

ISO 3166 is the International Standard for country codes and codes for their subdivisions. The purpose of ISO 3166 is to establish internationally recognised codes for the representation of names of countries, territories or areas of geographical interest, and their subdivisions.

(See them at http://www.iso.org/iso/country\_codes.htm and http://en.wikipedia.org/wiki/ISO\_ 3166-2)

Table A.4.1: ISO 3166-1 and ISO 3166-2 aliases				
Alias	Key	Description		
	GB-CAM			
	GB-ESS			
ALIAS:EAL	GB-HRT	East Anglia: United Kingdom region composed of the		
	GB-NFK	administrative counties of Norfolk to the north, Suffolk		
	GB-SFK	to the south, Cambridgeshire and Essex to the west.		

#### A.5 Institution ID

Please contact DCC if your institution is not listed, or you wish to modify the text

ID	Institution
001	Spanish National Cancer Research Centre (CNIO,
002	Madrid)  Barcelona Supercomputing Center (BSC-CNS,
002	Madrid)
003	EMBL-EBI (Hinxton)

## A.6 Value Codes for Platform or Validation Platform

Please contact the DCC if your platform/technology is not listed here.

Table	A.6.1: Value Codes for Platform or Validation Platform
Key	Platform or Validation Platform
1	PCR
2	qPCR
3	capillary sequencing
4	SOLiD sequencing
5	Illumina GA sequencing
6	454 sequencing
7	Helicos sequencing
8	Affymetrix Genome-Wide Human SNP Array 6.0
9	Affymetrix Genome-Wide Human SNP Array 5.0
10	Affymetrix Mapping 100K Array Set
11	Affymetrix Mapping 500K Array Set
12	Affymetrix Mapping 10K 2.0 Array Set
13	Affymetrix EMET Plus Premier Pack
14	Agilent Whole Human Genome Oligo Microarray Kit
15	Agilent Human Genome 244A
16	Agilent Human Genome 105A
17	Agilent Human CNV Association 2x105K
18	Agilent Human Genome 44K
19	Agilent Human CGH 1x1M
20	Agilent Human CGH 2x400K
21	Agilent Human CGH 4x180K
22	Agilent Human CGH 8x60K
23	Agilent Human CNV 2x400K
24	Agilent Human miRNA Microarray Kit (v2)
25	Agilent Human CpG Island Microarray Kit
26	Agilent Human Promoter ChIP-on-chip Microarray
27	Set
27	Agilent Human SpliceArray
28	Illumina human1m-duo
	Continued on next page

Table A.6.1 – continued from previous page

v3.0
/2.0D
nome
nome
e Set
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-07-2
-07-3
oaary
page

Table A.6.1 – concluded from previous page

Key	Platform or Validation Platform
67	Ion Torrent PGM
68	Illumina GoldenGate Methylation Cancer Panel I
69	Illumina Infinium HumanMethylation450
70	Agilent 8 x 15K Human miRNA-specific microarray
71	M.D. Anderson Reverse Phase Protein Array Core
<b>7</b> 2	Microsatellite Instability Analysis
73	Agilent 244K Custom Gene Expression G4502A-07
74	Illumina HumanCNV370-Duo v1.0 BeadChip
75	Illumina HumanOmniExpress BeadChip

## A.7 Chromosome Names for Reference Genome GRCh37

Table A.7.1: Chromosome Names for Reference Genome GRCh37

Key	Chromosome Name
1	1
2	2
3	3
4	4
5	5
6	6
7	7
8	8
9	9
10	10
11	11
12	12
13	13
14	14
15	15
16	16
17	17
18	18
19	19
20	20
21	21
22	22
23	X
24	Υ
25	MT
26	c5_H2
	Continued on next page

Table A.7.1 – continued from previous page

	Table A.7.1 – continued from previous page
Key	Chromosome Name
27	c6_COX
28	c6_QBL
29	NT_113870
30	NT_113871
31	NT_113872
32	 NT_113874
33	 NT_113878
34	 NT_113880
35	 NT_113881
36	 NT_113884
37	NT_113885
38	NT_113886
39	NT_113888
40	NT_113889
41	NT_113890
42	NT_113898
43	NT_113899
44	NT_113901
45	NT_113902
46	NT_113903
47	NT_113906
48	NT_113908
49	NT_113909
50	NT_113910
51	NT_113911
52	NT_113912
53	NT_113915
54	NT_113916
55	NT_113917
56	NT_113923
57	NT_113924
58	NT_113925
59	NT_113926
60	NT_113927
61	NT_113929
62	NT_113930
63	NT_113931
64	NT_113932
65	NT_113933
66	NT_113934
67	NT_113935
68	NT_113936
	Continued on next page

Table A.7.1 – continued from previous page

	Table A.7.1 – continued from previous page
Key	Chromosome Name
69	NT_113937
70	NT_113939
71	NT_113943
<b>7</b> 2	NT_113944
73	NT_113946
74	NT_113949
75	NT_113951
76	NT_113953
77	NT_113954
78	NT_113956
79	NT_113957
80	NT_113958
81	NT_113960
82	NT_113961
83	NT_113962
84	NT_113963
85	NT_113964
86	NT_113965
87	NT_113966
88	HSCHR17_1
89	HSCHR17_RANDOM_CTG2
90	HSCHR17_RANDOM_CTG3
91	HSCHR19_RANDOM_CTG2
92	HSCHR1_RANDOM_CTG12
93	HSCHR1_RANDOM_CTG5
94	HSCHR4_RANDOM_CTG2
95	HSCHR4_RANDOM_CTG3
96	HSCHR6_MHC_APD
97	HSCHR6_MHC_COX
98	HSCHR6_MHC_DBB
99	HSCHR6_MHC_MANN
100	HSCHR6_MHC_MCF
101	HSCHR6_MHC_QBL
102	HSCHR6_MHC_SSTO
103	HSCHR7_RANDOM_CTG1
104	HSCHR8_RANDOM_CTG1
105	HSCHR8_RANDOM_CTG4
106	HSCHR9_RANDOM_CTG2
107	HSCHR9_RANDOM_CTG4
108	HSCHR9_RANDOM_CTG5
109	HSCHRUN_RANDOM_CTG1
110	HSCHRUN_RANDOM_CTG10
	Continued on next page

Table A.7.1 – concluded from previous page

Key	Chromosome Name
111	HSCHRUN_RANDOM_CTG11
112	HSCHRUN_RANDOM_CTG13
113	HSCHRUN_RANDOM_CTG14
114	HSCHRUN_RANDOM_CTG15
115	HSCHRUN_RANDOM_CTG16
116	HSCHRUN_RANDOM_CTG17
117	HSCHRUN_RANDOM_CTG2
118	HSCHRUN_RANDOM_CTG20
119	HSCHRUN_RANDOM_CTG21
120	HSCHRUN_RANDOM_CTG22
121	HSCHRUN_RANDOM_CTG23
122	HSCHRUN_RANDOM_CTG26
123	HSCHRUN_RANDOM_CTG29
124	HSCHRUN_RANDOM_CTG3
125	HSCHRUN_RANDOM_CTG30
126	HSCHRUN_RANDOM_CTG31
127	HSCHRUN_RANDOM_CTG32
128	HSCHRUN_RANDOM_CTG33
129	HSCHRUN_RANDOM_CTG34
130	HSCHRUN_RANDOM_CTG35
131	HSCHRUN_RANDOM_CTG36
132	HSCHRUN_RANDOM_CTG4
133	HSCHRUN_RANDOM_CTG40
134	HSCHRUN_RANDOM_CTG5
135	HSCHRUN_RANDOM_CTG6
136	HSCHRUN_RANDOM_CTG9
137	HSCHR4_1

## A.8 Value Codes for Reference Genome Assembly Version

Table A.8.1: Value Codes for Reference Genome Assembly Version

Key	Reference Genome Assembly Version
1	GRCh37
2	NCBI36
3	GRCh37.p1
4	GRCh37.p2
5	GRCh37.p3
6	GRCh37.p4
7	GRCh37.p5

## A.9 Value Codes for Raw Data Repository

Table A.9.1: Value Codes for Raw Data Repository

Key Raw Data Repository

Continued on next page

Table A.9.1 – concluded from previous page

Key	Raw Data Repository
1	EGA
2	dbSNP
3	TCGA
4	CGHub
5	GE0