CRDC Submission Portal APIs

1. Introduction

This document walks you through the basics of submitting data to CRDC Submission Portal using APIs. Information provide in CRDC Data Submission Instructions will not be included in this document. It is recommended to get familiar with CRDC Data Submission Instructions and submission workflow using CRDC Data Submission Portal before interacting with CRDC Submission APIs.

CRDC Submission Portal provides a set of Graphql APIs (https://hub.datacommons.cancer.gov/api/graphql. Different functions are provided as separate API queries (read) and mutations (write). Users may use API tools like Postman as client to interact with the APIs. Users may also choose to write code in any popular programming languages to interact with the APIs.

All CRDC Submission Portal API queries and mutations use HTTP POST method. The APIs will always return HTTP code 200 (success) even when API call fails. The API returns data and errors in JSON format. If an API call succeeds, the returned data will be under "data" key. If any error occurs, the error information will be returned under "error" key ("data" key will be null).

CRDC Submission Portal API is authentication and authorization controlled. API tool (or your code) needs to be set up to send "Bearer Token" in "Authorization" header.

A Graphql API's schema contains definitions of all queries, mutations and their parameters and return types. This document will only provide information that cannot be found in the Graphql schema.

2. Prerequisites

- A valid API token downloaded from Data Submission Portal.
- API testing tool, such as Postman or GraphiQL or custom code.
- Knowledge of using API tools and/or interacting with APIs in code.
- Basic knowledge of Graphql APIs (https://graphql.org/).

3. Conventions

A dot separated property name is used to describe the hierarchy in the data. For example, **submissions**_id means_id property under **submissions** property. See screenshot below.

4. Starting a new submission

Before creating a data submission, user needs to determine which study the data submission will be submitted to. To retrieve approved studies for a user's organization, call API query **getMyUser**.

API signature:

```
getMyUser : User
```

Sample Query	Sample graphql variables
<pre>query getMyUser {</pre>	
getMyUser {	
_id	
studies {	
_id	
}	
}	
}	

Important return values:

• **studies._id**: should be used as **studyID** parameter of **createSubmission** API.

To create a new data submission, call API mutation **createSubmission**.

```
createSubmission (
    studyID: String!,
    dataCommons: String!,
    name: String!,
    intention: String!,
    dataType: String!
): Submission
```

```
Sample Query
                                                 Sample graphql variables
mutation createSubmission($studyID: String!,
$dataCommons: String!, $name:
                                                   "studyID": "4ab75167-0121-4fea-b515-
String!,$intention: String!, $dataType:
                                                 94c01d8380cc",
String!) {
                                                   "dataCommons": "ICDC",
 createSubmission(
                                                   "name": "API full workflow",
   studyID: $studyID
                                                   "intention": "New/Update",
   dataCommons: $dataCommons
                                                   "dataType": "Metadata and Data Files"
   name: $name
   intention: $intention
   dataType: $dataType
 ) {
   _id
   status
   createdAt
 }
}
```

- studyID: _id field of an approved study
- dataCommons: data common's name such as "CTDC", "ICDC" etc. For General Commons the name should be "CDS"
- **name**: a user selected name for the submission
- intention: should be one of ["New/Update", "Delete"]
- dataType: should be one of ["Metadata Only", "Metadata and Data Files"]

Important return values:

• _id: submission's ID, aka. submissionID

5. Continuing an existing submission

a. Retrieve a list of all submissions.

To retrieve a list of all submissions a user has access to, call API query listSubmissions.

```
listSubmissions(
   name: String,
   dbGaPID: String,
   dataCommons: String,
   submitterName: String,
   organization: String,
   status: [String],
   first: Int = -1,
```

```
offset: Int = 0,
orderBy: String = "updatedAt",
sortDirection: String = "DESC"): ListSubmissions
```

```
Sample Query
                                                  Sample graphql variables
query listSubmissions($first: Int, $offset:
Int, $orderBy: String, $sortDirection:
                                                    "status": "All",
String, $status: [String]) {
                                                    "first": 2,
 listSubmissions(
                                                    "offset": 0,
    first: $first
                                                    "sortDirection": "desc",
                                                    "orderBy": "updatedAt"
    offset: $offset
    orderBy: $orderBy
    sortDirection: $sortDirection
    status: $status
 } {
    total
    submissions {
      _id
      name
      submitterName
      organization {
        _id
        name
      }
      dataCommons
      studyAbbreviation
      dbGaPID
      modelVersion
      status
      conciergeName
      createdAt
      updatedAt
      intention
      dataFileSize {
        formatted
      }
  }
```

- Name: results will be filtered by data submission's name
- dbGaPID: results will be filtered by data submission's dbGaPID
- dataCommons: results will be filtered by data submission's data commons
- submitterName: results will be filtered by data submission's creator's name
- organization: results will be filtered by data submission's organization name,
 this parameter is deprecated and will be removed in the future

- **Status:** results will be filtered by this parameter if one or more of the following values is provided: ["New", "In Progress", "Submitted", "Released", "Completed", "Archived", "Canceled", "Rejected", "Withdrawn", "Deleted", "All"]. If "All" is provided, no filter will be applied.
- **first:** number of records to be returned, if -1 is sent, API will return all available data
- offset: skip given number of records before returning data
- orderBy: property name used to sort returned data
- sortDirection: should be one of ["ASC", "DESC"]

Important return values:

• **submissions._id:** submission's ID, aka. submissionID

b. Retrieve information about a submission

To retrieve detailed information about a submission, call API query **getSubmission**.

```
getSubmission(_id: ID!): Submission
```

```
Sample Query
                                                 Sample graphal variables
query getSubmission($id: ID!) {
                                                   "id": "eea2a531-4860-4e09-bf8e-151f73d4c379"
 getSubmission(_id: $id) {
   _id
   name
   submitterID
   submitterName
   organization {
      _id
      name
   dataCommons
   modelVersion
   studyID
   studyAbbreviation
   dbGaPID
   bucketName
   rootPath
   status
   metadataValidationStatus
   fileValidationStatus
   crossSubmissionStatus
   validationStarted
   validationEnded
```

```
validationScope
  validationType
  deletingData
  fileErrors {
    submissionID
    type
    validationType
    batchID
    displayID
    {\tt submittedID}
    severity
    uploadedDate
    validatedDate
    errors {
      title
      description
    warnings {
      title
      description
    }
  history {
    status
    reviewComment
    dateTime
    userID
  }
  conciergeName
  conciergeEmail
  intention
  dataType
  otherSubmissions
  createdAt
  updatedAt
}
```

• _id: submission's ID, aka. submissionID

To retrieve statistics of a submission, call API query **submissionStats**.

```
submissionStats(_id: ID!): SubmissionStats
```

• _id: submission's ID, aka. submissionID

To retrieve uploaded metadata, call API query getSubmissionNodes.

```
getSubmissionNodes(
   submissionID: String!,
   nodeType: String!,
   status: String = "All",
   nodeID: String,
   first: Int = 10,
   offset: Int = 0,
   orderBy: String = "nodeID",
   sortDirection: String = "ASC"
): SubmissionNodes
```

```
Sample Query
                                                 Sample graphql variables
query getSubmissionNodes($_id: String!,
$nodeType: String!, $status: String,
                                                   "_id": "eea2a531-4860-4e09-bf8e-
$submittedID: String, $first: Int, $offset:
                                                 151f73d4c379",
Int, $orderBy: String, $sortDirection:
                                                   "first": 20,
String) {
                                                    "offset": 0,
                                                    "sortDirection": "desc",
 getSubmissionNodes(
                                                   "orderBy": "studyID",
    submissionID: $_id
                                                    "nodeType": "study",
    nodeType: $nodeType
    status: $status
                                                   "status": "All",
                                                    "submittedID": ""
    nodeID: $submittedID
    first: $first
                                                 }
    offset: $offset
    orderBy: $orderBy
    sortDirection: $sortDirection
 ) {
    total
    IDPropName
    properties
    nodes {
     nodeID
        nodeType
        status
        props
 }
```

- submissionID: submission's ID
- **nodeType**: type of the metadata node to be returned
- **status:** should be one of ["All", "New", "Error", "Passed", "Warning"]. If "All" is provided. no filter will be applied, otherwise, return will be filtered by metadata's status.
- **nodeID:** if provided, return will be filtered by provided node ID, any node ID partially match given value will be returned.
- **first:** number of records to be returned, if -1 is sent, API will return all available data
- offset: skip given number of records before returning data
- orderBy: property name used to sort returned data
- **sortDirection:** should be one of ["ASC", "DESC"]

Important return values:

- **IDPropName**: name of the metadata node's ID property
- **Properties**: names of all metadata node's properties

• **Nodes**.props: a JSON string contains all properties of the metadata node, needs to be parsed as JSON in the code.

c. Uploading Files and Manifests

It is recommended to upload data files via Uploader CLI Tool. It is also possible to upload data files by writing code.

To retrieve an CLI configuration file, call API query **retrieveCLIConfig**. Returned data contains correct content and format (including line breaks and indentation). It is recommended to call this API in code and save returned string into a YAML file without modifying the content in anyway. For example, "my-cli-configuration.yml". API tools have their own way of displaying data contains line breaks, it will be hard to preserve original return data in an API tool.

API signature:

```
retrieveCLIConfig(
   submissionID: String!,
   apiURL: String!,
   dataFolder: String,
   manifest: String
): String
```

```
Sample QuerySample graphql variablesString, $manifest: String) {{retrieveCLIConfig("submissionID": "0997c282-d3ac-47ab-a7f2-2dacd2ca7d7c",submissionID: $submissionID"dataFolder": "/Users/user1/Desktop/datafiles/",apiURL: $apiURL"manifest": "/Users/user1/Desktop/datafiles/manifest.tsv",dataFolder: $dataFolder"apiURL": "https://hub-qa2.datacommons.cancer.gov/api/graphql"manifest: $manifest}
```

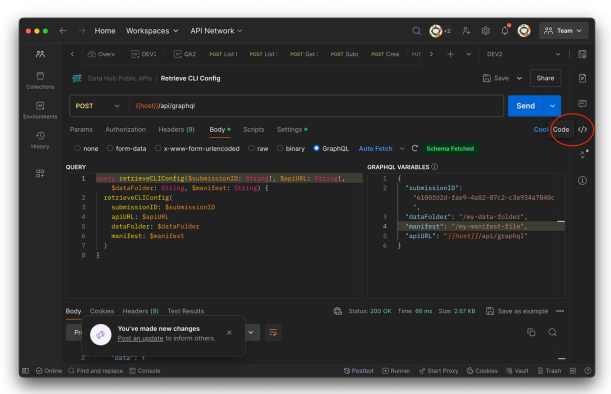
Important parameters:

- submissionID: submission's ID
- apiURL: must be "https://hub.datacommons.cancer.gov/api/graphql"
- dataFolder: local path of data files folder, for example "/Users/me/my-datafiles-folder"
- manifest: local path of manifest file, for example "/Users/me/my-metadata-folder/my-file-manifest.tsv"

To save CLI configuration file retrieved as a file without writing code, follow steps below:

- Setup API request in Postman
- Click "Code" button (</>) on the right-side bar

- Copy and paste the command into a terminal (cmd on Windows), then add following code to end of the command
 - " | jq -r '.["data"]["retrieveCLIConfig"]' > my-cli-configuration.yml"



If uploading data files in code is preferred, user may call API mutation **createTempCredentials** to retrieve a set of temporary credentials to use in the code.

API signature:

createTempCredentials (submissionID: ID!): TempCredentials

```
Sample QuerySample graphql variablesmutation createTempCredentials ($submissionID:{ID!) {"submissionID": "6100fd2d-fae9-4a82-87c2-c3e934a7840c"$submissionID) {}accessKeyIdsecretAccessKeysessionToken}}
```

Important parameters:

• submissionID: submission's ID

Metadata templates can be uploaded via Uploader CLI Tool. It is also possible to upload metadata templates via API. If using API is preferred, it is recommended to perform following steps in code rather than in API tool:

- Step 1: create a "batch" by calling API mutation createBatch.
- Step 2: upload metadata templates using pre-signed URLs retrieved in step 1.
- Step 3: update upload results by calling API mutation **updateBatch**. API will take the input as is and store the status of the files in the database. An asynchronous essential validation will be triggered by the API call, and the batch's status will be updated based on validation result. If validations passed, metadata will be loaded into submission database and status will be set to "**Uploaded**". Otherwise, batch status will be set to "**Failed**". Note, individual file status will be based on the input, and will not be updated by essential validation service.
- Step 4: retrieve essential validation results by calling API query listBatches

createBatch:

API signature:

```
createBatch (
  submissionID: ID!,
  type: String,
  files: [String!]!
): NewBatch
```

```
Sample Query
                                                  Sample graphql variables
mutation createBatch($submissionID: ID!,
$type: String, $files: [String!]!) {
                                                    "submissionID": "0997c282-d3ac-47ab-a7f2-
 createBatch(submissionID: $submissionID,
                                                  2dacd2ca7d7c",
type: $type, files: $files) {
                                                    "type": "metadata",
                                                    "files": ["program.tsv", "study.tsv",
    id
                                                  "participant.tsv", "sample.tsv"]
    submissionID
    bucketName
                                                  }
    filePrefix
    type
    fileCount
    files {
      fileName
      signedURL
    }
    status
    createdAt
    updatedAt
 }
```

Important parameters:

- submissionID: submission's ID
- type: should be one of ["metadata", "data file"]
- **files**: list of file names to be uploaded.

Important return values:

- _id: batch's internal ID, aka batchID
- **files.signedURL**: S3 pre-signed URL that can be used to upload metadata templates

updateBatch:

API signature:

```
updateBatch (batchID: ID!, files: [UploadResult]): Batch
```

```
Sample Query
                                                   Sample graphal variables
mutation updateBatch($batchID: ID!, $files:
[UploadResult]) {
                                                     "batchID": "13af4bee-bed8-42ce-8252-
 updateBatch(batchID: $batchID, files: $files)
                                                  eb5a756185b9",
                                                     "files": [
{
    _id
                                                    {
    submissionID
                                                         "fileName": "program.tsv",
                                                         "succeeded": true,
    type
    fileCount
                                                         "errors": null
    files {
                                                      }, {
     filePrefix
                                                         "fileName": "study.tsv",
                                                         "succeeded": true,
      fileName
                                                         "errors": null
      size
      status
                                                      }, {
                                                         "fileName": "participant.tsv",
      errors
                                                         "succeeded": true,
      createdAt
                                                         "errors": null
      updatedAt
    }
                                                      }, {
    status
                                                         "fileName": "sample.tsv",
    createdAt
                                                         "succeeded": true,
    updatedAt
                                                         "errors": null
 }
                                                      }
}
                                                     ]
                                                  }
```

Important parameters:

- batchID: batch's internal ID
- **files.skipped:** reserved for CLI use, should set to false

Important return values:

• _id: batch's internal ID, aka batchID

- displayID: batch's UIID
- **status**: current status of the batch. It will change after essential validation is finished (explained in previous sections). Please call **listBatches** to get the latest status.

To retrieve information about all batches, call API query listBatches.

```
listBatches(
  submissionID: ID!,
  first: Int = 10,
  offset: Int = 0,
  orderBy: String = "updatedAt",
  sortDirection: String = "DESC"
): ListBatches
```

```
Sample Query
                                                  Sample graphql variables
query listBatches($submissionID: ID!, $first:
Int, $offset: Int, $orderBy: String,
                                                    "submissionID": "0997c282-d3ac-47ab-a7f2-
$sortDirection: String) {
                                                  2dacd2ca7d7c",
 listBatches(
                                                    "first": 20,
    submissionID: $submissionID
                                                    "offset": 0,
    first: $first
                                                    "sortDirection": "desc",
    offset: $offset
                                                    "orderBy": "createdAt"
    orderBy: $orderBy
    sortDirection: $sortDirection
    total
    batches {
        _id
        displayID
        createdAt
        updatedAt
        submissionID
        type
        fileCount
        files {
            nodeType
            filePrefix
            fileName
            size
            status
            errors
            createdAt
            updatedAt
```

```
status
errors
}
}
```

- submissionID: submission's ID
- **first:** number of records to be returned, if -1 is sent, API will return all available data
- offset: skip given number of records before returning data
- orderBy: property name used to sort returned data
- **sortDirection:** should be one of ["ASC", "DESC"]

Important return values:

- batches._id: batch's internal ID, aka batchID
- batches.displayID: batch's UI ID
- **batches.files.nodeType:** node type contained in the metadata file. This value is only available when the metadata file can be successfully read by the essential validation service. Otherwise, it will be null.

d. Deleting data

To delete a metadata node, call API mutation deleteDataRecords.

```
deleteDataRecords(
   submissionID: String!,
   nodeType: String!,
   nodeIDs: [String!]
): DataValidation
```

```
Sample Query
                                                 Sample graphql variables
mutation deleteDataRecords($_id: String!,
$nodeType: String!, $nodeIds: [String!]) {
                                                   "_id": "26000952-13c6-4cf5-9ac5-
 deleteDataRecords(submissionID: $_id,
                                                 cf87b0c942f9",
nodeType: $nodeType, nodeIDs: $nodeIds) {
                                                   "nodeType": "sample",
   success
                                                   "nodeIds": [
                                                     "allval_Samp_Sep18_01"
   message
 }
                                                   ]
}
                                                 }
```

- submissionID: submission's ID
- **nodeType**: type of the metadata node to be deleted, or "**data file**" if deleting data files from S3 bucket is desired.
- **nodeIDs:** a list of node IDs, API will delete metadata node that matches provided node IDs.

Important return values:

• **success**: a Boolean value indicates if a deletion operation has been successfully initialized, the deletion will be performed asynchronously.

To delete a data file from submission bucket, call API mutation deleteDataRecords.

API signature:

```
deleteDataRecords(
   submissionID: String!,
   nodeType: String!,
   nodeIDs: [String!]
): DataValidation
```

```
Sample Query
                                                 Sample graphql variables
mutation deleteDataRecords($_id: String!,
$nodeType: String!, $nodeIds: [String!]) {
                                                   "_id": "26000952-13c6-4cf5-9ac5-
 deleteDataRecords(submissionID: $_id,
                                                 cf87b0c942f9",
nodeType: $nodeType, nodeIDs: $nodeIds) {
                                                   "nodeType": "data file",
                                                   "nodeIds": [
   success
   message
                                                     "41 batchtextfiles.txt"
 }
                                                   ٦
                                                 }
```

Important parameters:

- submissionID: submission's ID
- nodeType: must be "data file"
- **nodeIDs:** a list of file names to be deleted.

Important return values:

 success: a Boolean value indicates if a deletion operation has been successfully initialized, the deletion will be performed asynchronously.

e. Running validations

To validate uploaded data, call API mutation **validateSubmission**. API signature:

validateSubmission(
 _id: ID!,
 types: [String]
 scope: String
): DataValidation

```
Sample QuerySample graphql variablesmutation validateSubmission($_id: ID!, $types:{[String], $scope: String) {"_id": "0997c282-d3ac-47ab-a7f2-validateSubmission(_id: $_id, types: $types,2dacd2ca7d7c",scope: $scope) {"types": [success"metadata", "data file"}],}"scope": "All"
```

- _id: submission's ID, aka. submissionID
- **types:** any combination of following values: ["metadata". "data file"]
- scope: should be one of ["New", "All"]

Important return values:

• **success**: a Boolean value indicates if a validation has been successfully initialized, it has no relationship to the validation's result.

To retrieve aggregated validation issues, call API query aggregatedSubmissionQCResults.

```
aggregatedSubmissionQCResults(
    submissionID: ID!,
    severity: String = "all"
    first: Int = 20,
    offset: Int = 0,
    orderBy: String = "count"
    sortDirection: String = "DESC"): aggregatedQCResults
```

```
Sample Query
                                                 Sample graphql variables
query submissionQCResults($submissionID: ID!,
$severity: String, $first: Int, $offset: Int,
                                                   "submissionID": "6100fd2d-fae9-4a82-87c2-
$orderBy: String, $sortDirection: String) {
                                                 c3e934a7840c",
 aggregatedSubmissionQCResults(
                                                   "first": -1,
   submissionID: $submissionID
                                                   "offset": 0,
                                                   "sortDirection": "desc",
   severity: $severity
                                                   "orderBy": "displayID",
   first: $first
   offset: $offset
                                                   "severity": "All"
   orderBy: $orderBy
   sortDirection: $sortDirection
 } (
   total
   results {
       title
        severity
        count
        code
 }
}
```

- submissionID: submission's ID
- **severities**: should be one of ["all", "error", "warning"], return will be filtered by given issue severity. "all" means both errors and warnings.
- **first:** number of records to be returned, if -1 is sent, API will return all available data
- offset: skip given number of records before returning data
- orderBy: should be one of ["count", "title", "code", "severity"]
- **sortDirection:** should be one of ["ASC", "DESC"]

Important return values:

- title: title of the issue
- **severity**: either "Error" or "Warning"
- count: number of occurrences of the issue
- **code**: issue code that can be used in **submissionQCResults** API as **issueCode** parameter

To retrieve detailed validation issues, call API query **submissionQCResults**.

```
submissionQCResults(
   _id: ID!,
   nodeTypes: [String],
   batchIDs: [ID],
   severities: String,
   issueCode: String,

first: Int = 10,
   offset: Int = 0
   orderBy: String = "uploadedDate",
   sortDirection: String = "DESC"
): QCResults
```

```
Sample Query
                                                 Sample graphql variables
query submissionQCResults($id: ID!,
$nodeTypes: [String], $batchIDs: [ID],
                                                   "id": "0997c282-d3ac-47ab-a7f2-
$severities: String, $first: Int, $offset: Int,
                                                 2dacd2ca7d7c",
$orderBy: String, $sortDirection: String) {
                                                   "first": -1,
 submissionQCResults(
                                                   "offset": 0,
   _id: $id
                                                   "sortDirection": "desc",
   nodeTypes: $nodeTypes
                                                   "orderBy": "displayID",
   batchIDs: $batchIDs
                                                   "severities": "All"
   severities: $severities
   first: $first
   offset: $offset
   orderBy: $orderBy
   sortDirection: $sortDirection
 ) {
   total
   results {
      submissionID
      type
      validationType
      batchID
      displayID
      submittedID
      severity
      uploadedDate
      validatedDate
      errors {
       title
       description
      warnings {
       title
       description
      }
```

```
}
```

- _id: submission's ID, aka. submissionID
- **nodeTypes:** a list of metadata node types or "data file", return will be filtered by given metadata node types or file validation results if "data file" is given.
- batchIDs: a list of batches' internal IDs, return will be filtered by given batch internal IDs
- **severities**: should be one of ["All", "Error", "Warning"], return will be filtered by given issue severity. "All" means both errors and warnings.
- issueCode: results will be filtered by the issue code
- **first:** number of records to be returned, if -1 is sent, API will return all available data
- offset: skip given number of records before returning data
- orderBy: property name used to sort returned data
- sortDirection: should be one of ["ASC", "DESC"]

Important return values:

- validationType: either "metadata" or "data file"
- submittedID: a metadata node's ID, or file name of a data file

f. Submitting your Final Dataset

To submit a submission for review, call API mutation submissionAction.

```
submissionAction (
  submissionID: ID!,
  action: String!
  comment: String
): Submission
```

```
name
}
dataCommons
modelVersion
studyID
studyAbbreviation
dbGaPID
bucketName
rootPath
status
metadataValidationStatus
fileValidationStatus
crossSubmissionStatus
validationStarted
validationEnded
validationScope
validationType
deletingData
fileErrors {
    submissionID
    type
    validationType
    batchID
    displayID
    submittedID
    severity
    uploadedDate
    validatedDate
    errors {
        title
        description
    }
    warnings {
        title
        description
    }
}
history {
    status
    reviewComment
    dateTime
    userID
}
conciergeName
conciergeEmail
intention
dataType
```

```
otherSubmissions
createdAt
updatedAt
}
}
```

- submissionID: submission's ID
- action: should be "Submit" for this use case. Valid value includes ["Submit", "Withdraw", "Cancel"]