

# Package ‘Rga4gh’

November 7, 2016

**Type** Package

**Title** An Interface to the GA4GH API

**Version** 0.1.1

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**Description**

An Interface to the GA4GH API that allows users to easily GET responses and POST requests to GA4GH Servers. See <<http://ga4gh.org>> for more information about the GA4GH project.

**Depends** R (>= 3.1.0), httr

**Imports** jsonlite

**Suggests** magrittr

**License** GPL-2

**LazyData** TRUE

**RoxygenNote** 5.0.1

**NeedsCompilation** no

**Repository** CRAN

**Date/Publication** 2016-11-07 21:07:40

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ga4gh_client	<i>GA4GH Client</i>
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## Description

‘ga4gh\_client’ creates an object with class `ga4gh_client` that you can use to call the API specified by the arguments in the function. This is so you can query an API multiple times without having to worry about where it is every time. You can also set some defaults for the client to determine how responses from the client typically behave. Creating these objects also makes it easier to query multiple servers with similar requests.

## Usage

```
ga4gh_client(server, port = NULL, api_location = "/ga4gh", log_level = 0,
authentication_key = "", page_size = 10)
```

**Arguments**

server	The URL of the GA4GH server
port	The port number the GA4GH server is running on
api_location	A string to suffix to the server giving the location of the API on the server. By default this is "/ga4gh" i.e. the API is at http(s)://IP_ADDRESS/ga4gh
log_level	The amount of debugging information to log
authentication_key	The authentication key provided by the server after logging in
page_size	The default number of results for the client to return for paged responses

**Value**

An object with class ga4ghClient

**Examples**

```
## Create a client for the reference server at http://1kgenomes.ga4gh.org
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")

## Not run:
## Use this client as the first argument in the high-level API functions

## Search for datasets in this server
library(magrittr)
datasets <- ref_client %>% search_datasets() %>% content()

## or specify the api location in the search function
search_datasets("http://1kgenomes.ga4gh.org")

## End(Not run)
```

get_bio_sample	<i>GET a Bio Sample</i>
----------------	-------------------------

**Description**

GET a Bio Sample

**Usage**

```
get_bio_sample(client, bio_sample_id)
```

**Arguments**

client	A ga4gh_client object
bio_sample_id	The ID of the BioSample

## Examples

```
## Create a client
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")
## Not run:
library(magrittr)
## Retrieve the object with id 'id'
bio_sample <- ref_client %>% get_bio_sample("id")

## End(Not run)
```

**get\_call\_set**

*GET a Call Set*

## Description

GET a Call Set

## Usage

```
get_call_set(client, call_set_id)
```

## Arguments

client	A ga4gh_client object
call_set_id	The ID of the Call Set

## Examples

```
## Create a client
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")
## Not run:
library(magrittr)
## Retrieve the object with id 'id'
call_set <- ref_client %>% get_call_set("id")

## End(Not run)
```

---

get_dataset	<i>GET a Dataset</i>
-------------	----------------------

---

## Description

GET a Dataset

## Usage

```
get_dataset(client, dataset_id)
```

## Arguments

client	A ga4gh_client object
dataset_id	The ID of the Dataset

## Examples

```
## Create a client
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")
## Not run:
library(magrittr)
## Retrieve the object with id 'id'
dataset <- ref_client %>% get_dataset("id")

## End(Not run)
```

---

get_expression_level	<i>GET an Expression Level</i>
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---

## Description

GET an Expression Level

## Usage

```
get_expression_level(client, expression_level_id)
```

## Arguments

client	A ga4gh_client object
expression_level_id	The ID of the Expression Level

## Examples

```
## Create a client
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")
## Not run:
library(magrittr)
## Retrieve the object with id 'id'
exp_level <- ref_client %>% get_expression_level("id")

## End(Not run)
```

**get\_feature**

*GET a Feature*

## Description

GET a Feature

## Usage

```
get_feature(client, feature_id)
```

## Arguments

client	A ga4gh_client object
feature_id	The ID of the Feature

## Examples

```
## Create a client
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")
## Not run:
library(magrittr)
## Retrieve the object with id 'id'
feature <- ref_client %>% get_feature("id")

## End(Not run)
```

---

get_feature_set	<i>GET a Feature Set</i>
-----------------	--------------------------

---

### Description

GET a Feature Set

### Usage

```
get_feature_set(client, feature_set_id)
```

### Arguments

client	A ga4gh_client object
feature_set_id	The ID of the Feature Set

### Examples

```
## Create a client
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")
## Not run:
library(magrittr)
## Retrieve the object with id 'id'
feature_set <- ref_client %>% get_feature_set("id")

## End(Not run)
```

---

---

get_ga4gh	<i>GET an Object</i>
-----------	----------------------

---

### Description

GET an object from the GA4GH API

### Usage

```
get_ga4gh(object, ...)

## S3 method for class 'ga4ghClient'
get_ga4gh(object, operation, id, ...)

## Default S3 method:
get_ga4gh(object, operation, id, port = NULL, ...)

## S3 method for class 'list'
get_ga4gh(object, operation, id, ...)
```

**Arguments**

object	An object to GET from. I.e. A ga4ghClient object, a url for the server or a list of either.
...	Arguments to pass to other methods
operation	The name of the operation. Eg "datasets"
id	The ID of the Object
port	The port number

**Examples**

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")  
  
## Not run:  
  
## Use the client and get an object from the api  
ref_client %>% get_ga4gh("datasets", "id")  
  
## Or provide the API location as a string  
get_ga4gh("http://1kgenomes.ga4gh.org", "datasets", "id")  
  
## End(Not run)
```

**get\_individual**      *GET an Individual*

**Description**

GET an Individual

**Usage**

```
get_individual(client, individual_id)
```

**Arguments**

client	A ga4gh_client object
individual_id	The ID of the Individual

## Examples

```
## Create a client
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")
## Not run:
library(magrittr)
## Retrieve the object with id 'id'
individual <- ref_client %>% get_individual("id")

## End(Not run)
```

---

get\_read\_group

*GET a Read Group*

---

## Description

GET a Read Group

## Usage

```
get_read_group(client, read_group_id)
```

## Arguments

client	A ga4gh_client object
read_group_id	The ID of the Read Group

## Examples

```
## Create a client
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")
## Not run:
library(magrittr)
## Retrieve the object with id 'id'
read_group <- ref_client %>% get_read_group("id")

## End(Not run)
```

`get_read_group_set`      *GET a Read Group Set*

### Description

GET a Read Group Set

### Usage

```
get_read_group_set(client, read_group_set_id)
```

### Arguments

<code>client</code>	A <code>ga4gh_client</code> object
<code>read_group_set_id</code>	The ID of the Read Group Set

### Examples

```
## Create a client
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")
## Not run:
library(magrittr)
## Retrieve the object with id 'id'
read_group_set <- ref_client %>% get_read_group_set("id")

## End(Not run)
```

`get_reference_id`      *GET a Reference*

### Description

GET a Reference

### Usage

```
get_reference(client, reference_id)
```

### Arguments

<code>client</code>	A <code>ga4gh_client</code> object
<code>reference_id</code>	The ID of the Reference

## Examples

```
## Create a client
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")
## Not run:
library(magrittr)
## Retrieve the object with id 'id'
reference <- ref_client %>% get_reference("id")

## End(Not run)
```

---

get\_reference\_set      *GET a Reference Set*

---

## Description

GET a Reference Set

## Usage

```
get_reference_set(client, reference_set_id)
```

## Arguments

client	A ga4gh_client object
reference_set_id	The ID of the Reference Set

## Examples

```
## Create a client
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")
## Not run:
library(magrittr)
## Retrieve the object with id 'id'
reference_set <- ref_client %>% get_reference_set("id")

## End(Not run)
```

`get_rna_quantification`  
*GET an RNA Quantification*

### Description

GET an RNA Quantification

### Usage

```
get_rna_quantification(client, rna_quantification_id)
```

### Arguments

<code>client</code>	A <code>ga4gh_client</code> object
<code>rna_quantification_id</code>	The ID of the RNA Quantification

### Examples

```
## Create a client
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")
## Not run:
library(magrittr)
## Retrieve the object with id 'id'
rna_quantification <- ref_client %>% get_rna_quantification("id")

## End(Not run)
```

`get_rna_quantification_set`  
*GET an RNA Quantification Set*

### Description

GET an RNA Quantification Set

### Usage

```
get_rna_quantification_set(client, rna_quantification_set_id)
```

### Arguments

<code>client</code>	A <code>ga4gh_client</code> object
<code>rna_quantification_set_id</code>	The ID of the RNA Quantification Set

## Examples

```
## Create a client
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")
## Not run:
library(magrittr)
## Retrieve the object with id 'id'
rna_quantification_set <- ref_client %>% get_rna_quantification_set("id")

## End(Not run)
```

---

### get\_variant

*GET a Variant*

---

## Description

GET a Variant

## Usage

```
get_variant(client, variant_id)
```

## Arguments

client	A ga4gh_client object
variant_id	The ID of the Variant

## Examples

```
## Create a client
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")
## Not run:
library(magrittr)
## Retrieve the object with id 'id'
variant <- ref_client %>% get_variant("id")

## End(Not run)
```

---

`get_variant_annotation_set`  
*GET a Variant Annotation Set*

---

**Description**

GET a Variant Annotation Set

**Usage**

```
get_variant_annotation_set(client, variant_annotation_set_id)
```

**Arguments**

<code>client</code>	A <code>ga4gh_client</code> object
<code>variant_annotation_set_id</code>	The ID of the Variant Annotation Set

**Examples**

```
## Create a client
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")
## Not run:
library(magrittr)
## Retrieve the object with id 'id'
variant_annotation_set <- ref_client %>% get_variant_annotation_set("id")

## End(Not run)
```

---

`get_variant_set`      *GET a Variant Set*

---

**Description**

GET a Variant Set

**Usage**

```
get_variant_set(client, variant_set_id)
```

**Arguments**

<code>client</code>	A <code>ga4gh_client</code> object
<code>variant_set_id</code>	The ID of the Variant Set

## Examples

```
## Create a client
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")
## Not run:
library(magrittr)
## Retrieve the object with id 'id'
variant_set <- ref_client %>% get_variant_set("id")

## End(Not run)
```

post\_ga4gh

*POST to a GA4GH Server*

## Description

Use this to POST a request to an operation. The operations should simply be specified as a path in the API i.e. "/datasets/search" to search datasets. The body can be json or an R list. Use the `search_*` functions for a simpler interface.

## Usage

```
post_ga4gh(client, operation, body)
```

## Arguments

<code>client</code>	A <code>ga4gh_client</code> object
<code>operation</code>	The API operation to POST to as a string. Eg "/datasets/search"
<code>body</code>	The body of the POST either as JSON or a list (which will be converted to json by <code>httr</code> )

## Examples

```
## Create a client
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "", log_level = 1)

## A request body as a list
body_list <- list(pageSize = 2)

## Not run:
## Post to the /datasets/search operation
datasets_list <- ref_client %>% post_ga4gh("/datasets/search", body_list) %>% content
datasets_list

## End(Not run)
```

`search_bio_samples`     *Search for Bio Samples*

## Description

Search for Bio Samples

## Usage

```
search_bio_samples(client, dataset_id = NULL, individual_id = NULL,
page_size = NULL, page_token = NULL)
```

## Arguments

<code>client</code>	A <code>ga4gh_client</code> object
<code>dataset_id</code>	The ID of the Dataset to search within
<code>individual_id</code>	Return BioSamples for the provided Individual ID
<code>name</code>	Return BioSamples with the given name found by case-sensitive string matching.
<code>page_size</code>	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
<code>page_token</code>	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of <code>nextPageToken</code> from the previous response.

## Examples

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")

## Not run:
library(magrittr)
## Find a dataset to search in
datasets <- ref_client %>% search_datasets() %>% content()
d_id <- datasets$datasets[[1]]$id

## Search for bio samples in the dataset
bio_samples <- ref_client %>% search_bio_samples(d_id) %>% content()
bio_samples

## End(Not run)
```

---

search\_call\_sets      *Search for Call Sets*

---

## Description

Search for Call Sets

## Usage

```
search_call_sets(client, variant_set_id = NULL, bio_sample_id = NULL,
  name = NULL, page_size = NULL, page_token = NULL)
```

## Arguments

client	A ga4gh_client object
variant_set_id	The ID of the Variant Set to search within
bio_sample_id	Return only call sets generated from the provided BioSample ID.
name	Only return call sets with this name (case-sensitive, exact match).
page_size	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
page_token	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of nextPageToken from the previous response.

## Examples

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")

## Not run:
library(magrittr)
## Find a dataset to search in
datasets <- ref_client %>% search_datasets() %>% content()
d_id <- datasets$datasets[[1]]$id

## Find a variant set to search in
variant_sets <- ref_client %>% search_variant_sets(d_id) %>% content()
vs_id <- variant_sets$variantSets[[1]]$id

## Search for call sets in the variant set
call_sets <- ref_client %>% search_call_sets(vs_id) %>% content()
call_sets

## End(Not run)
```

---

search_datasets	<i>Search for Datasets</i>
-----------------	----------------------------

---

**Description**

Search for Datasets

**Usage**

```
search_datasets(client, page_size = NULL, page_token = NULL)
```

**Arguments**

client	A ga4gh_client object
page_size	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
page_token	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of nextPageToken from the previous response.

**Examples**

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")

## Not run:
## Use the client to search for datasets
library(magrittr)
datasets <- ref_client %>% search_datasets() %>% content()
datasets

## Or specify the API location within search_datasets
search_datasets("http://1kgenomes.ga4gh.org")

## End(Not run)
```

---



---

search_expression_levels	<i>Search for Expression Levels</i>
--------------------------	-------------------------------------

---

**Description**

Search for Expression Levels

## Usage

```
search_expression_levels(client, rna_quantification_id = NULL,  
  feature_ids = NULL, threshold = NULL, page_size = NULL,  
  page_token = NULL)
```

## Arguments

client	A ga4gh_client object
rna_quantification_id	The ID of the RNA Quantification to search within
feature_ids	A list of Feature IDs. Only return expressions with any of the specified feature_ids.
threshold	Number
page_size	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
page_token	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of nextPageToken from the previous response.

## Examples

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")  
  
## Not run:  
  
library(magrittr)  
## Find a dataset to search in  
datasets <- ref_client %>% search_datasets() %>% content()  
d_id <- datasets$datasets[[1]]$id  
  
## Find an rna quantification set to search in  
rna_quantification_sets <- ref_client %>%  
  search_rna_quantification_sets(d_id) %>% content()  
rqs_id <- rna_quantification_sets$rnaQuantificationSets[[1]]$id  
  
## Find an rna quantification to search in  
rna_quantifications <- ref_client %>%  
  search_rna_quantifications(rqs_id) %>% content()  
rq_id <- rna_quantifications$rnaQuantifications[[1]]$id  
  
## Search for expression levels in the rna quantification  
expression_levels <- ref_client %>%  
  search_expression_levels(rq_id) %>% content()  
expression_levels  
  
## End(Not run)
```

---

search_features	<i>Search for Features</i>
-----------------	----------------------------

---

**Description**

Search for Features

**Usage**

```
search_features(client, feature_set_id = NULL, parent_id = NULL,
  feature_types = NULL, reference_name = NULL, gene_symbol = NULL,
  name = NULL, start = NULL, end = NULL, page_size = NULL,
  page_token = NULL)
```

**Arguments**

<code>client</code>	A <code>ga4gh_client</code> object
<code>feature_set_id</code>	The ID of the Feature Set to search within. Either <code>feature_set_id</code> or <code>parent_id</code> must be non-null.
<code>parent_id</code>	Restricts the search to direct children of the given parent feature ID. Either <code>feature_set_id</code> or <code>parent_id</code> must be non-null.
<code>feature_types</code>	A list of feature types. If specified, this query matches only annotations whose <code>feature_type</code> matches one of the provided ontology terms.
<code>reference_name</code>	Only return features on the reference with this name
<code>gene_symbol</code>	Only return features with matching the provided gene symbol (case-sensitive, exact match).
<code>name</code>	Only returns features with this name (case-sensitive, exact match).
<code>start</code>	The beginning of the window (0-based, inclusive) for which overlapping features should be returned. Genomic positions are non-negative integers less than reference length.
<code>end</code>	The end of the window (0-based, exclusive) for which overlapping features should be returned.
<code>page_size</code>	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
<code>page_token</code>	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of <code>nextPageToken</code> from the previous response.

**Examples**

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")  
## Not run:
```

```
library(magrittr)
## Find a dataset to search in
datasets <- ref_client %>% search_datasets() %>% content()
d_id <- datasets$datasets[[1]]$id

## Find a feature set to search in
feature_sets <- ref_client %>% search_feature_sets(d_id) %>% content()
fs_id <- feature_sets$featureSets[[1]]$id

## Search the for features in the feature set
features <- ref_client %>% search_features(fs_id) %>% content()
features$features[[1]]

## End(Not run)
```

---

search\_feature\_phenotype\_associations  
*Search for Feature Phenotype Associations*

---

**Description**

Search for Feature Phenotype Associations

**Usage**

```
search_feature_phenotype_associations(client, ...)
```

**Arguments**

client	A ga4gh_client object
...	Other named arguments for the body of the POST request

---

search\_feature\_sets    *Search for Feature Sets*

---

**Description**

Search for Feature Sets

**Usage**

```
search_feature_sets(client, dataset_id = NULL, page_size = NULL,
page_token = NULL)
```

## Arguments

<code>client</code>	A <code>ga4gh_client</code> object
<code>dataset_id</code>	The ID of the Dataset to search within
<code>page_size</code>	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
<code>page_token</code>	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of nextPageToken from the previous response.

## Examples

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")  
  
## Not run:  
library(magrittr)  
## Find a dataset to search in  
datasets <- ref_client %>% search_datasets() %>% content()  
d_id <- datasets$datasets[[1]]$id  
  
## Search for feature sets in the dataset  
feature_sets <- ref_client %>% search_feature_sets(d_id) %>% content()  
feature_sets  
  
## End(Not run)
```

`search_ga4gh`

*Search*

## Description

Search the GA4GH API

## Usage

```
search_ga4gh(object, ...)  
  
## S3 method for class 'ga4ghClient'  
search_ga4gh(object, operation, ...)  
  
## Default S3 method:  
search_ga4gh(object, port = NULL, operation, ...)  
  
## S3 method for class 'list'  
search_ga4gh(object, operation, ...)
```

**Arguments**

object	An object to POST to. I.e. A <code>ga4ghClient</code> object, a url for the server or a list of either.
...	Named fields for the JSON body
operation	The name of an operation. Eg "variantsets"
port	The port number

---

search_individuals	<i>Search for Individuals</i>
--------------------	-------------------------------

---

**Description**

Search for Individuals

**Usage**

```
search_individuals(client, dataset_id = NULL, name = NULL,
page_size = NULL, page_token = NULL)
```

**Arguments**

client	A <code>ga4gh_client</code> object
dataset_id	The ID of the Dataset to search within
name	Return Individuals with the given name found by case-sensitive string matching.
page_size	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
page_token	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of <code>nextPageToken</code> from the previous response.

**Examples**

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")

## Not run:
library(magrittr)
## Find a dataset to search in
datasets <- ref_client %>% search_datasets() %>% content()
d_id <- datasets$datasets[[1]]$id

## Search for individuals in the dataset
individuals <- ref_client %>% search_individuals(d_id) %>% content()
individuals

## End(Not run)
```

`search_phenotypes`      *Search for Phenotypes*

## Description

Search for Phenotypes

## Usage

```
search_phenotypes(client, phenotype_association_set_id = NULL,
                  description = NULL, id = NULL, qualifiers = NULL, type = NULL,
                  age_of_onset = NULL, page_size = NULL, page_token = NULL)
```

## Arguments

<code>client</code>	A <code>ga4gh_client</code> object
<code>phenotype_association_set_id</code>	The ID of Phenotype Association Set to search within
<code>description</code>	String
<code>id</code>	String
<code>qualifiers</code>	A <code>data.frame</code> with columns <code>id</code> , <code>sourceName</code> , <code>sourceVersion</code> , <code>term</code> .
<code>type</code>	A named list with names <code>id</code> , <code>sourceName</code> , <code>sourceVersion</code> , <code>term</code> . Only return results that match this type
<code>age_of_onset</code>	A named list with names <code>id</code> , <code>sourceName</code> , <code>sourceVersion</code> , <code>term</code> . Only return results that match this <code>age_of_onset</code>
<code>page_size</code>	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
<code>page_token</code>	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of <code>nextPageToken</code> from the previous response.

## Examples

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")

## Not run:
library(magrittr)
## Find a dataset to search in
datasets <- ref_client %>% search_datasets() %>% content()
d_id <- datasets$datasets[[1]]$id

## Find a phenotype association set to search in
phenotype_association_sets <- ref_client %>%
  search_phenotype_association_sets(d_id) %>% content()
pas_id <- phenotype_association_sets$phenotypeAssociationSets[[1]]$id
```

```
## Search for phenotypes in the phenotype association set
phenotypes <- ref_client %>% search_phenotypes(pas_id) %>% content()
phenotypes$phenotypes[[1]]

## End(Not run)
```

**search\_phenotype\_association\_sets***Search for Phenotype Association Sets***Description**

Search for Phenotype Association Sets

**Usage**

```
search_phenotype_association_sets(client, dataset_id = NULL,
                                  page_size = NULL, page_token = NULL)
```

**Arguments**

<code>client</code>	A <code>ga4gh_client</code> object
<code>dataset_id</code>	The ID of the Dataset to search within
<code>page_size</code>	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
<code>page_token</code>	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of <code>nextPageToken</code> from the previous response.

**Examples**

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")

## Not run:
library(magrittr)
## Find a dataset to search in
datasets <- ref_client %>% search_datasets() %>% content()
d_id <- datasets$datasets[[1]]$id

## Search for phenotype association sets in the dataset
phenotype_association_sets <- ref_client %>%
  search_phenotype_association_sets(d_id) %>% content()
phenotype_association_sets
```

---

```
## End(Not run)
```

---

**search\_reads***Search for Reads***Description**

Search for Reads

**Usage**

```
search_reads(client, read_group_ids, reference_id = NULL, start = NULL,
             end = NULL, page_size = NULL, page_token = NULL)
```

**Arguments**

<code>client</code>	A <code>ga4gh_client</code> object
<code>read_group_ids</code>	A list of Read Group IDs. The Read Groups to search. At least one id must be specified.
<code>reference_id</code>	The reference to query. Leaving blank returns results from all references, including unmapped reads - this could be very large.
<code>start</code>	The start position (0-based) of this query. If a reference is specified, this defaults to 0. Genomic positions are non-negative integers less than reference length. Requests spanning the join of circular genomes are represented as two requests one on each side of the join (position 0).
<code>end</code>	The end position (0-based, exclusive) of this query. If a reference is specified, this defaults to the reference's length.
<code>page_size</code>	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
<code>page_token</code>	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of nextPageToken from the previous response.

**Examples**

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")

## Not run:
library(magrittr)
## Find a dataset to search in
datasets <- ref_client %>% search_datasets() %>% content()
d_id <- datasets$datasets[[1]]$id
```

```

## Find a read group to search in
read_group_sets <- ref_client %>% search_read_group_sets(d_id) %>% content()
rg_id <- read_group_sets$readGroupSets[[1]]$readGroups[[1]]$id

## Find a reference_id to map reads (unmapped reads not supported yet)
rs_id <- ref_client %>% search_reference_sets %>% content() %>%
  `$(referenceSets)` %>% `[[`(1) %>% `$(id)`
ref_id <- ref_client %>% search_references(rs_id) %>% content() %>%
  `$(references)` %>% `[[`(1) %>% `$(id)`

## Search for reads in the read group set
reads <- ref_client %>% search_reads(rg_id, reference_id = ref_id) %>% content()
reads

## End(Not run)

```

**search\_read\_group\_sets***Search for Read Group Sets***Description**

Search for Read Group Sets

**Usage**

```
search_read_group_sets(client, dataset_id = NULL, bio_sample_id = NULL,
                      name = NULL, page_size = NULL, page_token = NULL)
```

**Arguments**

<code>client</code>	A <code>ga4gh_client</code> object
<code>dataset_id</code>	The ID of the Dataset to search within
<code>bio_sample_id</code>	Specifying the id of a BioSample record will return only readgroups with the given <code>bio_sample_id</code> .
<code>name</code>	Only return read group sets with this name (case-sensitive, exact match).
<code>page_size</code>	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
<code>page_token</code>	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of <code>nextPageToken</code> from the previous response.

## Examples

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")  
  
## Not run:  
library(magrittr)  
## Find a dataset to search in  
datasets <- ref_client %>% search_datasets() %>% content()  
d_id <- datasets$datasets[[1]]$id  
  
## Search for read group sets in the dataset  
read_group_sets <- ref_client %>% search_read_group_sets(d_id) %>% content()  
read_group_sets  
  
## End(Not run)
```

**search\_references**      *Search for References*

## Description

Search for References

## Usage

```
search_references(client, reference_set_id = NULL, accession = NULL,  
                 md5checksum = NULL, page_size = NULL, page_token = NULL)
```

## Arguments

<code>client</code>	A <code>ga4gh_client</code> object
<code>reference_set_id</code>	The ID of the Reference Set to search within
<code>accession</code>	Return the References for which the accession matches this string (case-sensitive, exact match).
<code>md5checksum</code>	Return the References for which the md5checksum matches this string (case-sensitive, exact match).
<code>page_size</code>	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
<code>page_token</code>	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of nextPageToken from the previous response.

## Examples

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")  
  
## Not run:  
library(magrittr)  
## Find a reference set to search in  
reference_sets <- ref_client %>% search_reference_sets() %>% content()  
rs_id <- reference_sets$referenceSets[[1]]$id  
  
## Search for references in the reference set  
references <- ref_client %>% search_references(rs_id) %>% content()  
references  
  
## End(Not run)
```

## search\_reference\_bases

*Search for Reference Bases*

### Description

Search for Reference Bases

### Usage

```
search_reference_bases(client, reference_id = NULL, start = NULL,  
end = NULL, page_size = NULL, page_token = NULL)
```

### Arguments

client	A ga4gh_client object
reference_id	The ID of the Reference to search within
start	The start position (0-based) of this query. Defaults to 0. Genomic positions are non-negative integers less than reference length. Requests spanning the join of circular genomes are represented as two requests one on each side of the join (position 0).
end	The end position (0-based, exclusive) of this query. Defaults to the length of this Reference
page_size	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
page_token	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of nextPageToken from the previous response.

## Examples

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")

## Not run:
library(magrittr)
## Find a reference set to search in
reference_sets <- ref_client %>% search_reference_sets() %>% content()
rs_id <- reference_sets$referenceSets[[1]]$id

## Find a reference to search in
references <- ref_client %>% search_references(rs_id) %>% content()
ref_id <- references$references[[1]]$id

## Find a string of bases from 10000 -> 11000
reference_bases <- ref_client %>%
  search_reference_bases(ref_id, start = 10000, end = 11000) %>% content()
reference_bases

## End(Not run)
```

**search\_reference\_sets** *Search for Reference Sets*

## Description

Search for Reference Sets

## Usage

```
search_reference_sets(client, assembly_id = NULL, accession = NULL,
  md5checksum = NULL, page_size = NULL, page_token = NULL)
```

## Arguments

client	A ga4gh_client object
assembly_id	Return the Reference Sets for which the assembly_id matches this string (case-sensitive, exact match).
accession	Return the Reference Sets for which the accession matches this string (case-sensitive, exact match).
md5checksum	Return the Reference Sets for which the md5checksum matches this string (case-sensitive, exact match).
page_size	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
page_token	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of nextPageToken from the previous response.

## Examples

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")  
  
## Not run:  
library(magrittr)  
## Search for reference sets  
reference_sets <- ref_client %>% search_reference_sets() %>% content()  
reference_sets  
  
## End(Not run)
```

---

### search\_rna\_quantifications

*Search for RNA Quantifications*

---

## Description

Search for RNA Quantifications

## Usage

```
search_rna_quantifications(client, rna_quantification_set_id,  
                           page_size = NULL, page_token = NULL)
```

## Arguments

client	A ga4gh_client object
rna_quantification_set_id	The ID of the RNA Quantification Set to search within
page_size	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
page_token	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of nextPageToken from the previous response.

## Examples

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")  
  
## Not run:  
  
library(magrittr)  
## Find a dataset to search in  
datasets <- ref_client %>% search_datasets() %>% content()  
d_id <- datasets$datasets[[1]]$id
```

```

## Find an rna quantification set to search in
rna_quantification_sets <- ref_client %>%
  search_rna_quantification_sets(d_id) %>% content()
rqs_id <- rna_quantification_sets$rnaQuantificationSets[[1]]$id

## Search for rna quantifications in the rna quantification set
rna_quantifications <- ref_client %>%
  search_rna_quantifications(rqs_id) %>% content()
rna_quantifications

## End(Not run)

```

**search\_rna\_quantification\_sets***Search for RNA Quantification Sets***Description**

Search for RNA Quantification Sets

**Usage**

```
search_rna_quantification_sets(client, dataset_id = NULL, page_size = NULL,
                               page_token = NULL)
```

**Arguments**

<code>client</code>	A <code>ga4gh_client</code> object
<code>dataset_id</code>	The ID of the Dataset to search within
<code>page_size</code>	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
<code>page_token</code>	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of <code>nextPageToken</code> from the previous response.

**Examples**

```

ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")

## Not run:

library(magrittr)
## Find a dataset to search in
datasets <- ref_client %>% search_datasets() %>% content()
d_id <- datasets$datasets[[1]]$id

```

```
## Search for rna quantification sets in the dataset
rna_quantification_sets <- ref_client %>%
  search_rna_quantification_sets(d_id) %>% content()
rna_quantification_sets

## End(Not run)
```

---

search_variants	<i>Search for Variants</i>
-----------------	----------------------------

---

## Description

Search for Variants

## Usage

```
search_variants(client, variant_set_id, call_set_ids = NULL,
  reference_name = NULL, start = NULL, end = NULL, page_size = NULL,
  page_token = NULL)
```

## Arguments

client	A ga4gh_client object
variant_set_id	The ID of the Variant Set to search within
call_set_ids	A list of Call Set IDs. Only return variant calls which belong to call sets with these IDs. If unspecified, return all variants and no variant call objects.
reference_name	Only return variants on this reference.
start	The beginning of the window (0-based, inclusive) for which overlapping variants should be returned. Genomic positions are non-negative integers less than reference length.
end	The end of the window (0-based, exclusive) for which overlapping variants should be returned.
page_size	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
page_token	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of nextPageToken from the previous response.

## Examples

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")  
  
## Not run:  
library(magrittr)  
## Find a dataset to search in  
datasets <- ref_client %>% search_datasets() %>% content()  
d_id <- datasets$datasets[[1]]$id  
  
## Find a variant set to search in  
variant_sets <- ref_client %>% search_variant_sets(d_id) %>% content()  
vs_id <- variant_sets$variantSets[[1]]$id  
  
## Search for variants in the variant set  
variants <- ref_client %>% search_variants(vs_id, reference_name = "1") %>%  
  content()  
variants$variants[[1]]  
  
## End(Not run)
```

## search\_variant\_annotations

*Search for Variant Annotations*

### Description

Search for Variant Annotations

### Usage

```
search_variant_annotations(client, variant_annotation_set_id,  
  reference_id = NULL, reference_name = NULL, start = NULL, end = NULL,  
  effects = NULL, page_size = NULL, page_token = NULL)
```

### Arguments

client	A ga4gh_client object
variant_annotation_set_id	The ID of the Variant Annotation Set to search within.
reference_id	Only return variants with reference alleles on the reference with this ID. One of this field or referenceName is required.
reference_name	Only return variants with reference alleles on the reference with this name. One of this field or referenceId is required.

start	The beginning of the window (0-based, inclusive) for which variants with overlapping reference alleles should be returned. Genomic positions are non-negative integers less than reference length. Requests spanning the join of circular genomes are represented as two requests one on each side of the join (position 0).
end	The end of the window (0-based, exclusive) for which variants with overlapping reference alleles should be returned.
effects	A data.frame with columns id, sourceName, sourceVersion, term. This filter allows variant, transcript combinations to be extracted by effect type(s). Only return variant annotations including any of these effects and only return transcript effects including any of these effects. Exact matching across all fields of the Sequence Ontology OntologyTerm is required. (A transcript effect may have multiple SO effects which will all be reported.) If empty, return all variant annotations.
page_size	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
page_token	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of nextPageToken from the previous response.

## Examples

```

ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")

## Not run:
## Find a dataset to search in
datasets <- ref_client %>% search_datasets() %>% content
d_id <- datasets$datasets[[1]]$id

## Find a variant set to search in
variant_sets <- ref_client %>% search_variant_sets(d_id) %>% content()
vs_id <- variant_sets$variantSets[[2]]$id

## Find a variant annotation set to search in
variant_annotation_sets <- ref_client %>%
  search_variant_annotation_sets(vs_id) %>% content()
vas_id <- variant_annotation_sets$variantAnnotationSets[[1]]$id

## Search for variant annotations in the variant annotation set
variant_annotations <- ref_client %>%
  search_variant_annotations(vas_id, reference_name = "1") %>% content
variant_annotations$variantAnnotations[[1]]

## End(Not run)

```

---

**search\_variant\_annotation\_sets**  
*Search for Variant Annotation Sets*

---

**Description**

Search for Variant Annotation Sets

**Usage**

```
search_variant_annotation_sets(client, variant_set_id, page_size = NULL,
                               page_token = NULL)
```

**Arguments**

client	A ga4gh_client object
variant_set_id	The ID of the Variant Set to Search
page_size	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
page_token	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of nextPageToken from the previous response.

**Examples**

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")

## Not run:
## Find a dataset to search in
datasets <- ref_client %>% search_datasets() %>% content
d_id <- datasets$datasets[[1]]$id

## Find a variant set to search in
variant_sets <- ref_client %>% search_variant_sets(d_id) %>% content()
vs_id <- variant_sets$variantSets[[2]]$id

## Search for variant annotation sets in the variant set
variant_annotation_sets <- ref_client %>%
  search_variant_annotation_sets(vs_id) %>% content()
variant_annotation_sets

## End(Not run)
```

---

**search\_variant\_sets** *Search for Variant Sets*

---

**Description**

Search for Variant Sets

**Usage**

```
search_variant_sets(client, dataset_id, page_size = NULL, page_token = NULL)
```

**Arguments**

client	A ga4gh_client object
dataset_id	The ID of the Dataset to search within
page_size	Specifies the maximum number of results to return in a single page. If unspecified, the client default will be used.
page_token	The continuation token, which is used to page through large result sets. To get the next page of results, set this parameter to the value of nextPageToken from the previous response.

**Examples**

```
ref_client <- ga4gh_client("http://1kgenomes.ga4gh.org", api_location = "")  
  
## Not run:  
library(magrittr)  
## Find a dataset to search in  
datasets <- ref_client %>% search_datasets() %>% content()  
d_id <- datasets$datasets[[1]]$id  
  
## Search for variantsets in the dataset  
variant_sets <- ref_client %>% search_variant_sets(d_id) %>% content()  
variant_sets  
  
## End(Not run)
```

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