

Package: Rega (via r-universe)

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Title R Interface to European Genome-Phenome Archive

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Description The European Genome-phenome Archive (EGA) provides long-term storage and controlled sharing of personally identifiable genetic data. The Rega package offers a streamlined and extensible R interface to the EGA API, facilitating the programmatic upload of metadata. GEO-like Excel submission template is provided as a default method of organizing submission metadata.

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BugReports <https://github.com/ivanek/Rega/issues>

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add_required_str	<i>Mark Required Fields with a Prefix</i>
------------------	---

Description

Mark Required Fields with a Prefix

Usage

```
add_required_str(p, r, req_str = "* ")
```

Arguments

p	Character vector. All fields to be processed.
r	Character vector. Fields that are required.
req_str	Character. Prefix to mark required fields. Defaults to "* ".

Value

A character vector with required fields prefixed and ordered to appear before non-required fields.

Examples

```
# Mark required fields with a prefix
add_required_str(c("Name", "Id", "Age"), c("Id", "Name"))
```

aliases_formatter *Format Aliases from a Table*

Description

Format Aliases from a Table

Usage

```
aliases_formatter(tab, params)
```

Arguments

tab	Data frame. The input table where the first row contains column names.
params	List. Additional parameters for formatting. Takes a formatter params value from parser parameter yaml file. Currently unused.

Value

A named list where each name corresponds to a formatted column name, and values are non-NA elements of the respective column.

Examples

```
tab <- data.frame(Alias = c("name1", "name2", NA), Value = c(1, 2, 3))
aliases_formatter(tab, params = list())
```

api_function_factory *Generate an API Function from Operation and Specification*

Description

This function dynamically creates an API function based on a given operation definition and API specification. The generated function handles URL construction, parameter validation, request execution, and response parsing.

Usage

```
api_function_factory(  
  op,  
  api,  
  verbosity = 0,  
  bearer_token = NULL,  
  token_url = .EGA_TOKEN_URL  
)
```

Arguments

op	List. The API operation definition, including method, path, parameters, and request body schema.
api	List. The API specification, including host and global security definitions.
verbosity	Integer, optional, values 0-3. Indicates with which verbosity level should the requests <code>httr2::req_perform</code> be performed. Default: 0.
bearer_token	Character, optional. The API bearer token for authentication, will be included in the headers of the request. Defaults to NULL
token_url	Character, optional. Token endpoint URL from which to obtain the access token. If <code>bearer_token</code> is specified, it will take precedence. Defaults to <code>.EGA_TOKEN_URL = "https://idp.ega-archive.org/realms/EGA/protocol/openid-connect/token"</code> .

Value

A dynamically generated function that performs the specified API operation. The function accepts arguments corresponding to operation parameters and executes the request using `httr2`.

Examples

```
api <- extract_api()
opdefs <- extract_operation_definitions(api)

# Generate an API function for a specific operation
f <- api_function_factory(
  opdefs[["get_files"]], api,
  bearer_token = "my_key"
)

# Call the generated function with parameters (requires credentials)
try(
  result <- f(status = "value1", prefix = "value2")
)
```

api_name_to_label *Convert API Names to Prettified Labels*

Description

This function converts API-style names with underscores into human-readable labels by replacing underscores with spaces and applying title case.

Usage

```
api_name_to_label(x)
```

Arguments

x Character vector. API field names to be converted.

Value

A character vector with API names converted to human-readable labels.

Examples

```
api_name_to_label(c("first_name", "last_name", "instument_model"))
```

column_table_formatter

Format a Column Table

Description

Format a Column Table

Usage

```
column_table_formatter(tab, params)
```

Arguments

tab Data frame. The input table submission metadata file where the first row contains column names.

params List. Additional parameters for formatting. Takes a formatter params value from parser parameter yaml file. Currently unused.

Value

A cleaned data frame with column names set from the first row, empty rows removed, and whitespace trimmed from all values.

Examples

```
df <- data.frame(
  ...1 = c("* Alias", "Sample1", "Sample2"),
  ...2 = c("* Phenotype", "wt", "ko"),
  ...3 = c("Description", NA, NA)
)

column_table_formatter(df, list())
```

create_client	<i>Generate API Client Functions</i>
---------------	--------------------------------------

Description

This function creates a named list of functions for interacting with an API, based on its specification and operation definitions.

Usage

```
create_client(api, ...)
```

Arguments

api	List. The API specification, including operation definitions, host, and global settings.
...	List. List of additional arguments passed to <code>api_function_factory</code> .

Value

A named list of functions, where each function corresponds to an API operation. The function names match the operation IDs from the specification.

Examples

```
client <- create_client(  
  extract_api(),  
  bearer_token = "my_key", verbosity = 1  
)  
  
# Call an operation using the client (requires credentials)  
try(  
  result <- client$get__files(status = "value1", prefix = "value2")  
)
```

default_parser	<i>Parser for a Default EGA Excel Template</i>
----------------	--

Description

This function parses the `extdata/ega_full_template_v3.xlsx` using the bundled parser parameter file in `extdata/default_parser_params.yaml` to extract information for EGA submission into format that can be easily passed into EGA API endpoints.

Usage

```
default_parser(metadata_file, param_file = NULL)
```

Arguments

`metadata_file` Character. Path to a default template xlsx file containing the submission metadata information.

`param_file` Character. Path to a yaml file with parameters for parser. If NULL, uses the `extdata/default_parser_params.yaml`. Defaults to NULL.

Value

List of data frames or lists. Submission information parsed from the xlsx file.

Examples

```
default_parser(
  system.file("extdata/submission_example.xlsx", package = "Rega")
)
```

default_validator *Validator for Default Parser*

Description

Used to validate internal consistency of submission metadata parsed using the default parser. Performs several checks on EGA dataset for submission, ensuring that aliases for studies, experiments, samples, runs, analyses and datasets are properly linked, as they will be replaced with provisional or accession IDs during submission process. Displays a success message if all validation passed or a summary message if validation failed. In addition it returns a data frame with validation details.

Usage

```
default_validator(meta, aliases = NULL)
```

Arguments

`meta` List of data frames. Correspond to tables of EGA submission.

`aliases` List of lists. Aliases that should present in the EGA tables. If NULL, the function will attempt to locate it in the `meta` parameter. Defaults to NULL

Value

Data frame. Validator object that includes all performed validations and their statistics (number of passes, fails and NAs, or whether errors or warnings were encountered during validation)

Examples

```
minimal_metadata <- list(
  aliases = list(
    studies = "Study1", experiments = "Experiment1",
    datasets = "Dataset1", samples = "Sample1", runs = "Run1",
    analyses = "Analysis1"
  ),
  files = tibble::tibble(
    file = "raw.fastq.gz", ega_file = list("raw.fastq.gz.c4gh")
  ),
  submission = tibble::tibble(title = "Submission"),
  studies = tibble::tibble(
    study = "Study1", title = "Study Title",
    description = "Study Description",
    study_type = "Whole Genome Sequencing"
  ),
  samples = tibble::tibble(
    alias = "Sample1", phenotype = "wild-type",
    biological_sex = "female", subject_id = "ID1"
  ),
  experiments = tibble::tibble(
    study = "Study1", experiment = "Experiment1",
    design_description = "Experiment Design",
    library_selection = "RANDOM", instrument_model_id = 1L,
    library_layout = "SINGLE", library_strategy = "WGS",
    library_source = "GENOMIC"
  ),
  runs = tibble::tibble(
    run = "Run1", experiment = "Experiment1", run_file_type = "srf",
    alias = "Sample1", files = list("raw.fastq.gz.c4gh")
  ),
  datasets = tibble::tibble(
    dataset = "Dataset1", title = "Dataset Title",
    description = "Dataset Description",
    policy_accession_id = "EGAP0000000001",
    dataset_types = list("Whole genome sequencing"),
    runs = list("Run1")
  )
)

default_validator(minimal_metadata)
```

delete_submission

Delete a Submission and Log Responses

Description

Deletes a submission identified by its ID using the client and logs the response if a logfile is specified.

Usage

```
delete_submission(id, client = NULL, logfile = NULL, ...)
```

Arguments

id	A string representing the submission identifier (provisional ID).
client	An API client object with a delete method for submissions.
logfile	A string specifying the path to a log file. If NULL, no log is written. Defaults to NULL.
...	Additional arguments for future extensions (currently unused).

Value

A list containing the response for the submission deletion.

Examples

```
mock_client <- list(  
  delete_submissions__provisional_id =  
    function(id) list(status = "deleted")  
)  
delete_submission("5678901", mock_client)
```

delete_submission_contents

Delete Submission Contents and Log Responses

Description

Deletes all data associated with a submission ID using the client and logs the responses if a logfile is specified.

Usage

```
delete_submission_contents(id, client = NULL, logfile = NULL, ...)
```

Arguments

id	A string representing the submission identifier. Can be either an accession or provisional ID.
client	List of functions. EGA API client created by create_client function from EGA API schema with delete methods. If NULL, default client will be created by create_client(extract_api()). Defaults to NULL.
logfile	A string specifying the path to a log file. If NULL, no log is written. Defaults to NULL.
...	Additional arguments for future extensions (currently unused).

Value

A list of responses for the deletion of associated datasets, analyses, runs, experiments, samples, and studies.

Examples

```
mock_client <- list(
  "delete__submissions__provisional_id__datasets" =
    function(id) list(status = "deleted")
)
delete_submission_contents(5678901, client = mock_client)
```

DELIM_CONVERTERS

Default Column Converters for EGA Metadata

Description

A named list of functions used to coerce delimited strings into specific data types during metadata processing.

Usage

```
DELIM_CONVERTERS
```

Format

A named list:

pubmed_ids Coerces values to integer.

Examples

```
df <- list(pub_sheet = data.frame(
  pubmed_ids = c(" 123456, 789", NA, "1001 ")
))

process_delimited_column(
  df, "pubmed_ids", separator = ",", converters = DELIM_CONVERTERS
)
```

`ega_oauth`*Set The OAUTH With EGA Username And Password*

Description

`ega_oauth` implements the EGA OAuth resource owner password flow, as defined by Section 4.3 of RFC 6749. It allows the user to supply their password once, exchanging it for an access token that can be cached locally. Please avoid entering the password directly when calling this function as it will be captured by `.Rhistory`.

Usage

```
ega_oauth(  
  req,  
  username = .get_ega_username(),  
  password = .get_ega_password(),  
  token_url = .EGA_TOKEN_URL  
)
```

Arguments

<code>req</code>	A <code>httr2</code> request.
<code>username</code>	Character. EGA User name. Defaults to the value returned by <code>.get_ega_username()</code> .
<code>password</code>	Character. EGA user Password. Defaults to the value returned by <code>.get_ega_password()</code> .
<code>token_url</code>	Character. The URL for the EGA token endpoint. Defaults to <code>.EGA_TOKEN_URL</code> = "https://idp.ega-archive.org/realms/EGA/protocol/openid-connect/token".

Value

returns a modified HTTP request that will use OAuth

Examples

```
req <- httr2::request("https://example.com/")  
  
# Request OAuth with default credentials  
try(oauth_req <- ega_oauth(req))  
  
# Request OAuth with custom credentials  
oauth_req <- ega_oauth(req, username = "user", password = "pass")
```

ega_token	<i>Retrieve EGA API Bearer Token</i>
-----------	--------------------------------------

Description

This function retrieves an API token from the European Genome-Phenome Archive (EGA) using user credentials.

Usage

```
ega_token(  
  username = .get_ega_username(),  
  password = .get_ega_password(),  
  token_url = .EGA_TOKEN_URL  
)
```

Arguments

username	Character. The username for EGA authentication. Defaults to the value returned by <code>.get_ega_username()</code> .
password	Character. The password for EGA authentication. Defaults to the value returned by <code>.get_ega_password()</code> .
token_url	Character. The URL for the EGA token endpoint. Defaults to the standard EGA token URL if not provided. Defaults to <code>.EGA_TOKEN_URL = "https://idp.ega-archive.org/realM"</code>

Value

A list containing the token details if successful. Actual token value can be retrieved by `token$access_token`

Examples

```
try(  
  ega_token(username = "my_username", password = "my_password")  
)  
  
try(  
  ega_token(token_url = "https://www.example.com")  
)
```

extract_api

Extract API Specification and Host Details

Description

This function parses an API specification file (JSON or YAML) and extracts relevant details.

Usage

```
extract_api(spec_file = NULL, host = NULL)
```

Arguments

spec_file	Character. Optional. Path to the API specification file in JSON or YAML format. If NULL default <code>extdata/ega_api_deref.yaml</code> is used. Defaults to NULL
host	Character. Optional. The API host URL. If not supplied, it will be inferred from the specification file's <code>servers</code> element. Defaults to NULL

Value

A list containing the parsed API specification, including the host and `basePath` elements. If the specification file lacks required elements, appropriate warnings or errors are raised.

Examples

```
# Extract API details from a default YAML specification file
api <- extract_api()

# Extract API details with a custom host
api <- extract_api(host = "https://api.example.com")
```

extract_operation_definitions*Extract API Operation Definitions*

Description

This function extracts operation definitions from an API specification, including HTTP methods, paths, parameters, request bodies, and responses.

Usage

```
extract_operation_definitions(api)
```

Arguments

`api` List. Parsed API specification, generated from a JSON or YAML file. Must include a `paths` element with API endpoint definitions.

Value

A named list of operations, where each name corresponds to an operation ID. If operation ID is not found in the specification, unique one will be created. Each operation contains:

- `method`: HTTP method (e.g., GET, POST).
- `path`: Endpoint path.
- `parameters`: List of operation parameters.
- `requestBody`: Details of the request body (if any).
- `responses`: Possible responses for the operation.
- `security`: Security requirements for the operation.

Examples

```
# Extract operation definitions from a parsed API specification
opdefs <- extract_operation_definitions(extract_api())
opdefs[["post__submissions"]]
```

`extract_resource_name` *Extract Resource Name from API Response URL*

Description

Extracts the specific resource identifier (e.g., "users", "datasets") from the path of an `httr2` response object by parsing the segment immediately following `/api/`.

Usage

```
extract_resource_name(resp)
```

Arguments

`resp` An `httr2_response` object.

Value

A character string containing the resource name.

Examples

```
resp <- httr2::response(  
  method = "GET",  
  url = "https://www.example.com/api/files"  
)  
extract_resource_name(resp)
```

fetch_files

Fetch file information from EGA inbox

Description

Query the remote client for requested file prefix, test whether a file is found for every element of `file_list` and return the server response.

Usage

```
fetch_files(file_list, client = NULL)
```

Arguments

`file_list` A character vector or list of file prefixes to check.

`client` List of functions. EGA API client created by `create_client` function from EGA API schema. If `NULL`, default client will be created by `create_client(extract_api())`. Defaults to `NULL`.

Value

Data frame. Parsed response from client API for requested files.

Examples

```
mock_client <- list(  
  get__files = function(prefix = NULL) {  
    data.frame(provisional_id = 12345, ega_relative_path = prefix)  
  }  
)  
fetch_files(c("file_a", "file_b"), mock_client)
```

file_formatter	<i>Format File Table with EGA File Paths</i>
----------------	--

Description

Format File Table with EGA File Paths

Usage

```
file_formatter(tab, params)
```

Arguments

tab	Data frame. The input table containing file information. Columns file, ega_inbox_relative_path need to be present in the data.
params	List. Additional parameters for formatting. Takes a formatter params value from parser parameter yaml file. Includes crypt_ext for encryption file extensions and prepend_slash to control path prefix.

Value

A formatted data frame with cleaned column names, and updated ega_file paths based on file and relative path information.

Examples

```
params <- list(prefix = "", crypt_ext = "c4gh", prepend_slash = FALSE)

# Dummy data, first row will be moved to column names
tab <- data.frame(
  x1 = c("file", "value1", "value2"),
  x2 = c("ega_inbox_relative_path", NA, "proj1")
)

file_formatter(tab, params)
```

filter_id_fields	<i>Filter Out ID Fields from a Character Vector</i>
------------------	---

Description

This function filters out elements from a character vector that match a specified regular expression pattern, removing ID fields.

Usage

```
filter_id_fields(x, pattern = NULL)
```

Arguments

x	Character vector. The input vector to filter.
pattern	Character. Optional. A regular expression pattern for matching ID fields to exclude. Defaults to "(?!policy_)accession_id provisional_id", which removes accession IDs and provisional IDs, but not policy accession ID.

Value

A character vector with elements matching the pattern removed.

Examples

```
# Filter out default ID fields
fields <- c("accession_id", "policy_accession_id", "name", "provisional_id")
filter_id_fields(fields)

# Filter with a custom pattern
filter_id_fields(fields, pattern = "_id")
```

finalise_submission *Finalise an EGA submission*

Description

Submits the finalisation request for a submission identified by either an accession or provisional ID. Validates the release date and sends optional dataset changelogs.

Usage

```
finalise_submission(
  id,
  release_date,
  dataset_changelogs = data.frame(),
  client = NULL,
  logfile = NULL,
  ...
)
```

Arguments

id	Character scalar. The submission accession or provisional ID.
release_date	Character scalar. Expected release date in YYYY-MM-DD format.
dataset_changelogs	Data frame. Optional changelog metadata for associated datasets. If specified, the required columns are dataset and message. Defaults to empty data.frame.
client	List of functions. EGA API client created by create_client function from EGA API schema. If NULL, default client will be created by create_client(extract_api()). Defaults to NULL.
logfile	Character. Path of log file to log the httr2 responses from individual operations or NULL. Defaults to NULL.
...	List. Additional arguments to the function.

Value

The API response object from the finalisation request.

Examples

```
# Requires credentials
try(
  finalise_submission("123456", "2025-12-31")
)
```

first_row_to_colnames *Use First Row as Column Names for a Data Frame*

Description

Use First Row as Column Names for a Data Frame

Usage

```
first_row_to_colnames(df, to_api = TRUE)
```

Arguments

df	Data frame. The input data frame whose first row will become column names.
to_api	Logical. Whether to convert labels to API-style names using label_to_api_name(). Defaults to TRUE.

Value

A data frame with updated column names and the first row removed.

Examples

```
df <- data.frame(id = c("A B", "C D_"), value = c("* E F", "GH"))

first_row_to_colnames(df)
first_row_to_colnames(df, to_api = FALSE)
```

fold_column	<i>Fold Columns with a Common Prefix into a Single Column Nested as List</i>
-------------	--

Description

If NA values are present in any of the columns to be nested, they will be removed. If the column is not present it will be added with NA as a single value.

Usage

```
fold_column(tab, column_prefix, new_name)
```

Arguments

tab	Data frame. The input table with columns to fold.
column_prefix	Character. The prefix of columns to nest into a single column represented as list.
new_name	Character. The name of the new folded column.

Value

A data frame with the specified columns nested into a single column.

Examples

```
tab <- data.frame(id = c(1, 2), name.1 = c("A1", NA), name.2 = c("B1", "B2"))
fold_column(tab, "name", "folded_column")
```

format_chromosomes *Format Chromosome Metadata*

Description

Formats and processes chromosome-related metadata from an input object by applying chromosome group lookups or splitting chromosome strings from the EGA enums.

Usage

```
format_chromosomes(metadata)
```

Arguments

metadata List. A list of data frames representing metadata sheets, containing analyses. Each row in analyses can have entry in chromosomes or chromosome_groups column.

Value

A list of formatted chromosome data extracted or computed from the input metadata.

Examples

```
# Mock metadata data frame
metadata <- list(
  analyses = data.frame(
    chromosomes = I(list(
      NA,
      list("group1--1--chr1--name1", "group2--3--chr3--name3"),
      "group1--2--chr2--name2"
    )),
    chromosome_groups = c("group1", NA, "group3"),
    stringsAsFactors = FALSE
  ),
  select_input_data = list(
    chromosomes = c("group1--1--chr1--name1", "group1--2--chr2--name2")
  )
)

format_chromosomes(metadata)
```

get_chr_group *Retrieve Chromosome Belonging to a Group*

Description

Retrieve Chromosome Belonging to a Group

Usage

```
get_chr_group(group_id, chr_enum, sep = "--")
```

Arguments

group_id	Character. The group ID to filter by.
chr_enum	Character vector. Chromosome enumeration data, where each element is a string containing fields separated by sep. Enum data is created by parse_enum and get_enum functions. See vignette for more details.
sep	Character. Field separator in a string. Defaults to "--"

Value

An integer vector of chromosome IDs corresponding to the specified group ID.

Examples

```
get_chr_group(
  "group1", c("group1--1--chr1--name1", "group2--2--chr2--name2")
)
```

get_entry_by_title *Retrieve EGA entries by title*

Description

Searches for entries across specified EGA metadata types that match a given title string. Returns a list of data frames for each type.

Usage

```
get_entry_by_title(title, type = NULL, client = NULL, logfile = NULL, ...)
```

Arguments

title	Character scalar. The title or substring to search for.
type	Character vector. One or more metadata types ("submissions", "studies", "samples", "experiments", "runs", "analyses" and "datasets"). If NULL, searches all valid types.
client	List of functions. EGA API client created by create_client function from EGA API schema. If NULL, default client will be created by create_client(extract_api()). Defaults to NULL.
logfile	Character. Path of log file to log the httr2 responses from individual operations or NULL. Defaults to NULL.
...	List. Additional arguments to the function.

Value

A named list of data frames containing entries matching the title.

Examples

```
# Requires credentials
try(
  get_entry_by_title("My Study", type = "studies")
)
```

get_enum

Retrieve Enum Values from an API

Description

This function retrieves the values of a specified enum from an API by invoking the corresponding client function.

Usage

```
get_enum(client, enum_name, enum_prefix = "get__enums__")
```

Arguments

client	List. The API client, typically generated by create_client, containing functions for API operations.
enum_name	Character. The name of the enum to retrieve.
enum_prefix	Character. Optional. The prefix used in the client for enum functions. Defaults to "get__enums__".

Value

The values of the specified enum.

Examples

```
# Create API client with mock api_key
client <- create_client(extract_api(), token_url = "ABCD")

# Retrieve enum values from the API client (requires credentials to work)
try(
  platform_models <- get_enum(client, enum_name = "platform_models")
)
```

get_formatter

Retrieve a Formatter Function by Type of Submission Metadata Table

Description

Retrieve a Formatter Function by Type of Submission Metadata Table

Usage

```
get_formatter(x, params)
```

Arguments

x Character. The name of the submission metadata table/sheet.
params List. A list containing a formatter element from parser parameter yaml file.

Value

The formatter function corresponding to the specified table.

Examples

```
# Load formatter params
params <- yaml::read_yaml(system.file(
  "extdata/default_parser_params.yaml",
  package = "Rega"
))

# Dummy data, first row will be moved to column names
tab <- data.frame(
  x1 = c("file", "value1", "value2"),
  x2 = c("ega_inbox_relative_path", NA, "proj1")
)

ff <- get_formatter("files", params)
ff_params <- get_formatter_params("files", params)

ff(tab, ff_params)
```

get_formatter_params *Retrieve Formatter Parameters by Name*

Description

Retrieve Formatter Parameters by Name

Usage

```
get_formatter_params(x, params)
```

Arguments

x Character. The name of the formatter for which to retrieve parameters.

params List. A list containing a formatter element from parser parameter yaml file.

Value

A list of parameters for the specified formatter.

Examples

```
# Load formatter params
params <- yaml::read_yaml(system.file(
  "extdata/default_parser_params.yaml",
  package = "Rega"
))

# Dummy data, first row will be moved to column names
tab <- data.frame(
  x1 = c("file", "value1", "value2"),
  x2 = c("ega_inbox_relative_path", NA, "proj1")
)

ff <- get_formatter("files", params)
ff_params <- get_formatter_params("files", params)

ff(tab, ff_params)
```

get_operation_schema *Retrieve the Schema for an API Operation*

Description

Retrieve the Schema for an API Operation

Usage

```
get_operation_schema(op)
```

Arguments

op List. The API operation definition containing a requestBody element with content and schema details.

Value

The schema for the operation's JSON request body, or NULL if no schema is defined.

Examples

```
# Get operations from API
opdefs <- extract_operation_definitions(extract_api())

# Retrieve the schema for a specific operation
schema <- get_operation_schema(opdefs[["post__submissions"]])
```

get_or_post *Retrieve or Submit Data to an EGA API Endpoint*

Description

This function retrieves existing data from an API or submits new data if it does not exist, with optional error handling and retrieval options.

- If no data is present in the database, supplied data will be inserted.
- If there is data already present in the database and the number of records don't match, error will be raised.
- If the number of records match and retrieve is set to TRUE data will be retrieved from database and nothing will be inserted. If retrieve is set to FALSE, error will be raised.

Usage

```
get_or_post(  
  submission_id,  
  data,  
  client,  
  endpoint,  
  retrieve = FALSE,  
  id_type = "provisional"  
)
```

Arguments

submission_id	An integer representing the submission provisional ID.
data	A data frame to be submitted.
client	An API client object with get and post methods.
endpoint	A string specifying the EGA API endpoint. The endpoint will be a submission type endpoint identified with provisional ID.
retrieve	A logical flag indicating whether to retrieve data if it already exists. Defaults to FALSE.
id_type	A string specifying type of EGA id. One of 'provisional' or 'accession'. Defaults to provisional.

Value

A data frame containing the response from the API.

Examples

```
# Create mock client for API endpoint  
mock_client <- list(  
  get__submissions__provisional_id__endpoint = function(id) {  
    message("Mock GET request")  
    # Simulate an empty response (no existing data)  
    return(NULL)  
  },  
  post__submissions__provisional_id__endpoint = function(id, body) {  
    message("Mock POST request")  
    message(body) # Simulate returning submitted data  
  }  
)  
  
# Create mock data to test the function  
test_data <- data.frame(id = 1:3, value = c("A", "B", "C"))  
  
# Test the function with mock data and client  
result <- get_or_post(  
  submission_id = 12345,  
  data = test_data,  
  client = mock_client,
```

```
    endpoint = "endpoint",  
    retrieve = FALSE  
  )
```

`get_properties`*Extract and Format Properties from a Schema*

Description

This function extracts property names from a schema, optionally filters out ID fields, and applies formatting such as marking required fields and prettifying the labels.

Usage

```
get_properties(schema, filter_ids = TRUE)
```

Arguments

`schema` List. The schema containing properties and required fields.
`filter_ids` Logical. Whether to filter out ID fields from the properties. Defaults to TRUE.

Value

A character vector of formatted property names and indications of required fields.

Examples

```
schemas <- get_schemas(extract_api())  
  
# Extract and format properties from a schema  
get_properties(schemas[[5]])  
  
# Extract properties without filtering ID fields  
get_properties(schemas[[6]], filter_ids = FALSE)
```

get_schemas	<i>Retrieve Request schemas from an API Specification</i>
-------------	---

Description

This function extracts and returns schemas related to requests from the API specification.

Usage

```
get_schemas(api)
```

Arguments

api	List. The API specification, typically containing a components element with nested schemas.
-----	---

Value

A list of schemas whose names contain "Request", filtered from the schemas element of the API specification.

Examples

```
# Extract request schemas from an API specification
request_schemas <- get_schemas(extract_api())
```

get_sentence_number	<i>Count the number of sentences in text</i>
---------------------	--

Description

Count how many sentences are in a character string, based on terminal punctuation marks ., !, or ? following an alphanumeric character. If there is no punctuation character at the end, it will still count it as another sentence.

Usage

```
get_sentence_number(text)
```

Arguments

text	A character vector.
------	---------------------

Value

An integer scalar giving the number of sentences in text.

Examples

```
get_sentence_number("First sentence. Second sentence? Third!")
```

get_submission	<i>Retrieve Submission Data and Log Responses</i>
----------------	---

Description

Retrieves data associated with a submission ID using the client and logs the responses if a logfile is specified.

Usage

```
get_submission(id, client = NULL, logfile = NULL, ...)
```

Arguments

id	A string representing the submission identifier. Can be either an accession or provisional ID.
client	List of functions. EGA API client created by <code>create_client</code> function from EGA API schema with get methods. If NULL, default client will be created by <code>create_client(extract_api())</code> . Defaults to NULL.
logfile	A string specifying the path to a log file. If NULL, no log is written. Defaults to NULL.
...	Additional arguments for future extensions (currently unused).

Value

A list of responses including submission data and associated datasets, analyses, runs, experiments, samples, and studies.

Examples

```
mock_client <- list(
  "get__submissions__accession_id" = function(id) list(data = id),
  "get__submissions__accession_id__datasets" =
    function(id) list(datasets = id)
)
get_submission("EGAB12345678901", mock_client)
```

get_word_number	<i>Count the number of words in text</i>
-----------------	--

Description

Compute the number of words in each element of a character vector using non-word separators.

Usage

```
get_word_number(text)
```

Arguments

text A character vector with text.

Value

An integer vector giving the number of words per element of text.

Examples

```
get_word_number(c("one two", "three four five"))
```

has_linked_sheets	<i>Check for Linked Sheets in Metadata</i>
-------------------	--

Description

Determines whether a specified sheet is present and contains at least one non-NA value in the provided metadata.

Usage

```
has_linked_sheets(metadata, colname)
```

Arguments

metadata A list of data frame objects to check.
colname A string specifying the name of the column to look for.

Value

A logical vector indicating whether each element of metadata contains the specified column with at least one non-NA value.

Examples

```
metadata <- list(
  sheet1 = list(sheet_name = c(1, NA)),
  sheet2 = list(other_name = NA)
)
has_linked_sheets(metadata, "sheet_name")
```

is_accession

Check if a String is a Valid Accession Identifier.

Description

Verifies whether the input string matches the format of a valid accession identifier based on a specified schema.

Usage

```
is_accession(x, schema = NULL)
```

Arguments

x	A character vector to be tested for validity as accessions.
schema	A character string specifying the schema. Valid options include "study", "studies", "sample", "samples", "experiment", "experiments", "analysis", "analyses", "run", "runs", "policy", "DAC", "dataset", "datasets", "submission" and NULL. NULL will check against any of the schemas. Defaults to NULL.

Value

A logical vector indicating which values are accession IDs.

Examples

```
is_accession("EGAB0000000001", "submission") # TRUE
is_accession("EGA12345678901", "sample") # FALSE
```

is_provisional	<i>Check whether IDs is a provisional ID.</i>
----------------	---

Description

Determine if input values match the format of provisional IDs, either as whole-number numerics or as character strings of at least two digits without leading zeros.

Usage

```
is_provisional(x)
```

Arguments

x A numeric or character vector of candidate provisional IDs.

Value

A logical vector indicating which values are provisional IDs.

Examples

```
is_provisional(c(10, 11, 3.5, 9))
```

is_valid_http_method	<i>Validate HTTP Method</i>
----------------------	-----------------------------

Description

Checks whether a given HTTP method is valid based on a predefined list of accepted methods (matches on lowercase).

Usage

```
is_valid_http_method(m)
```

Arguments

m A string representing the HTTP method to validate.

Value

A logical value: TRUE if m is a valid HTTP method, otherwise FALSE.

Examples

```
is_valid_http_method("GET") # TRUE
is_valid_http_method("get") # TRUE
is_valid_http_method("DELETE") # TRUE
is_valid_http_method("foo") # FALSE
is_valid_http_method(NULL) # FALSE
```

label_to_api_name *Convert Prettified Labels to API Names*

Description

Convert Prettified Labels to API Names

Usage

```
label_to_api_name(x, req_str = "* ")
```

Arguments

x Character vector. Prettified labels to convert.

req_str Character. Optional prefix to remove from labels. Defaults to "* ".

Value

A character vector with labels converted to API-style names.

Examples

```
label_to_api_name(c("* First Name", "Last Name"))
label_to_api_name(c("# Instrument Model", "# Fragment SD"), req_str = "# ")
```

link_sheet *Link Data Between Metadata Sheets*

Description

Data frames representing metadata sheets that contain column names corresponding to sheet_name containing an ID reference (to a first column in sheet_name) will be replaced with the rest of the values nested as a list.

Usage

```
link_sheet(metadata, sheet_name)
```

Arguments

metadata List. A list of data frames representing metadata sheets.
 sheet_name Character. The name of the sheet to link with other sheets.

Value

A list of updated metadata with the specified values replaced based on referenced values present in sheet_name.

Examples

```
# Link data from a specific sheet to other sheets in metadata
metadata <- list(
  sheet1 = data.frame(id = c(1, 2), linked_sheet = c("A", "B")),
  linked_sheet = data.frame(id = c("A", "B"), value = c(10, 20))
)
updated_metadata <- link_sheet(metadata, "linked_sheet")
```

lut_add

*Add a Column to a Data Frame Based on Lookup Table***Description**

This function adds a new column to a data frame by mapping values from an existing column through a lookup table.

Usage

```
lut_add(df, to, from, lut)
```

Arguments

df A data frame to which the new column will be added.
 to A string specifying the name of the new column.
 from A string specifying the name of the column to map values from.
 lut A named list or vector serving as the lookup table.

Value

The input data frame with the added column.

Examples

```
df <- data.frame(id = c("A", "B", "C"), stringsAsFactors = FALSE)
lut <- list(A = 1, B = 2, C = 3)
lut_add(df, "value", "id", lut)
```

merge_linked_sheet	<i>Merge Linked Sheets with Source Data</i>
--------------------	---

Description

Merges a target column with a source column in a linked sheet's data, processing it into a format suitable for JSON parsing. Includes API-specific adjustments for certain data.

Usage

```
merge_linked_sheet(target, source, dat, sheet)
```

Arguments

target	A vector containing the target values.
source	A string specifying the source column to merge on.
dat	A data frame representing the data to be linked.
sheet	A string specifying the name of the sheet, used for API-specific processing.

Value

A data frame containing the merged data, or an empty list if the target is entirely NA.

Examples

```
target <- c(1, 2, 3)
source <- "id"
dat <- data.frame(id = c(1, 2, 3), value = c("A", "B", "C"))
merge_linked_sheet(target, source, dat, "collaborators")
```

multi_lut_add	<i>Add Multiple Lookup-Based Columns to a Data Frame</i>
---------------	--

Description

Adds multiple columns to a data frame by applying multiple lookup tables, each defined by a set of arguments specifying the new column, the source column, and the lookup table.

Usage

```
multi_lut_add(df, ...)
```

Arguments

`df` A data frame to which new columns will be added.

`...` A series of lists, each containing three elements: the name of the new column (`to`), the name of the source column (`from`), and the lookup table (`lut`). See [lut_add](#) for details.

Value

The input data frame with the added columns.

Examples

```
df <- data.frame(id = c("A", "B", "C"), stringsAsFactors = FALSE)
lut1 <- list(A = 1, B = 2, C = 3)
lut2 <- list(A = "x", B = "y", C = "z")
multi_lut_add(df, list("value1", "id", lut1), list("value2", "id", lut2))
```

na_to_empty_list *Convert NA Values to Empty Lists*

Description

Replaces NA values in a list with empty lists, preserving the original structure of the list. Doesn't work on nested lists.

Usage

```
na_to_empty_list(l)
```

Arguments

`l` A list containing elements that may include NA values.

Value

A list where any NA values have been replaced with empty lists.

Examples

```
input_list <- list(1, NA, "text", NA)
na_to_empty_list(input_list)
```

 new_submission

Submit New Data to EGA

Description

This function creates a new submission and associates all specified data with it. Following data has to be present in the request data object: submission studies, experiments, samples, runs, analyses, datasets. The files associated with the submission must be present in the EGA Inbox and they are fetched and matched according to Inbox path. In case the submission is interrupted or fails, all the information entered into EGA database is rolled back apart from the submission itself. If the workflow successfully creates a submission, but fails in the following steps, the returned submission ID can be used as a parameter to the workflow to continue entering data into existing submission. If logfile is specified, the responses from successfully executed steps (even if the error occurs), will be saved.

Usage

```
new_submission(
  dat,
  client = NULL,
  logfile = NULL,
  submission_id = NULL,
  retrieve = FALSE,
  ...
)
```

Arguments

dat	List of data frames. Parsed submission metadata containing correctly formatted and linked information for submission
client	List of functions. EGA API client created by <code>create_client</code> function from EGA API schema. If NULL, default client will be created by <code>create_client(extract_api())</code> . Defaults to NULL.
logfile	Character. Path of log file to log the <code>httr2</code> responses from individual operations or NULL. Defaults to NULL.
submission_id	Integer.
retrieve	Logical.
...	List. Additional arguments to the function.

Value

List of data frames. Parsed response objects from `httr2` requests

Examples

```

minimal_metadata <- list(
  aliases = list(
    studies = "Study1", experiments = "Experiment1",
    datasets = "Dataset1", samples = "Sample1", runs = "Run1",
    analyses = "Analysis1"
  ),
  files = tibble::tibble(
    file = "raw.fastq.gz", ega_file = list("raw.fastq.gz.c4gh")
  ),
  submission = tibble::tibble(title = "Submission"),
  studies = tibble::tibble(
    study = "Study1", title = "Study Title",
    description = "Study Description",
    study_type = "Whole Genome Sequencing"
  ),
  samples = tibble::tibble(
    alias = "Sample1", phenotype = "wild-type",
    biological_sex = "female", subject_id = "ID1"
  ),
  experiments = tibble::tibble(
    study = "Study1", experiment = "Experiment1",
    design_description = "Experiment Design",
    library_selection = "RANDOM", instrument_model_id = 1L,
    library_layout = "SINGLE", library_strategy = "WGS",
    library_source = "GENOMIC"
  ),
  runs = tibble::tibble(
    run = "Run1", experiment = "Experiment1", run_file_type = "srf",
    alias = "Sample1", files = list("raw.fastq.gz.c4gh")
  ),
  datasets = tibble::tibble(
    dataset = "Dataset1", title = "Dataset Title",
    description = "Dataset Description",
    policy_accession_id = "EGAP0000000001",
    dataset_types = list("Whole genome sequencing"),
    runs = list("Run1")
  )
)

ega <- create_client(extract_api(), verbosity = 0)

# Requires credentials
try(
  new_submission(minimal_metadata, ega)
)

```

Description

Parses the body of a body of httr2 response object from the EGA API, handling JSON and plain text content, and formats it into a tibble for further processing.

Usage

```
parse_ega_body(resp)
```

Arguments

resp An HTTP response object from the EGA API.

Value

A tibble containing the parsed and formatted response data. If the response is plain text without a JSON-like structure, a one-column tibble is returned with the raw content.

Examples

```
# Example with JSON response
json_resp <- httr2::response(
  method = "GET",
  url = "https://www.example.com/api/files",
  status = 200,
  headers = list("content-type" = "application/json"),
  body = charToRaw('{"id": 1, "name": "test"}')
)
parse_ega_body(json_resp)

# Example with plain text response
text_resp <- httr2::response(
  method = "POST",
  url = "https://www.example.com/api/submissions",
  status = 200,
  headers = list("content-type" = "text/plain"),
  body = charToRaw("Sample response text")
)
parse_ega_body(text_resp)
```

parse_enum

Parse Enum into a Formatted String

Description

This function parses an enum, represented as a data frame or character vector, into a formatted string for display or further use.

Usage

```
parse_enum(enum, sep = "--")
```

Arguments

enum	Data frame or character vector. The enum to parse. If a data frame, its rows are concatenated into strings. If a character vector, its elements are joined with newlines.
sep	Character. If enum has multiple fields, they will be pasted into a single string using this separator. Defaults to --

Value

A single string representing the parsed enum. Rows are joined by newlines and multiple enum fields are joined by sep.

Examples

```
# Parse an enum as a data frame
df_enum <- data.frame(key = c("A", "B"), value = c("1", "2"))
parse_enum(df_enum)

# Parse an enum as a character vector
vec_enum <- c("A", "B", "C")
parse_enum(vec_enum)
```

parse_json_body	<i>Parse and Standardize JSON Response Body</i>
-----------------	---

Description

Extracts the JSON body from a response and ensures the output is structured as a list of objects. Named lists (single records) are wrapped in a parent list to maintain consistency for downstream unnesting.

Usage

```
parse_json_body(resp)
```

Arguments

resp	An httr2_response object containing JSON content.
------	---

Value

A list of lists, where each inner list represents a record.

Examples

```
json_resp <- httr2::response(  
  method = "GET",  
  url = "https://www.example.com/api/files",  
  status = 200,  
  headers = list("content-type" = "application/json"),  
  body = charToRaw('[{ "id": 1, "name": "test" }]')  
)  
parse_json_body(json_resp)
```

parse_text_body

Parse Plain Text or JSON-like Response Body

Description

Processes a text response by either parsing it as JSON (if structured with curly braces or square brackets) or returning it as a list. Null JSON elements are converted to empty lists to facilitate unnesting.

Usage

```
parse_text_body(resp)
```

Arguments

resp An httr2_response object with "text/plain" content.

Value

A list of parsed data or a tibble if the content is raw text.

Examples

```
text_resp <- httr2::response(  
  method = "POST",  
  url = "https://www.example.com/api/submissions",  
  status = 200,  
  headers = list("content-type" = "text/plain"),  
  body = charToRaw("Sample response text")  
)  
parse_text_body(text_resp)
```

process_chromosomes *Process a Vector or List of Chromosome Data*

Description

This function processes chromosome data by extracting unique chromosome IDs and labels or retrieving chromosome group information from a lookup where applicable.

Usage

```
process_chromosomes(chr_data, select_input_data)
```

Arguments

`chr_data` A list containing chromosome-related information. Expected to have items chromosomes (scalar, vector or list) and/or chromosome_groups (scalar).

`select_input_data` A list containing look-up data for chromosomes.

Value

A data frame with chromosome id and label if chromosomes are present. If only chromosome groups exist, returns the result of lookup against the `select_input_data` with `get_chr_group()`. If neither are present, returns an empty list.

Examples

```
select_input_data <- list(
  chromosomes = c("group1--1--chr1--name1", "group1--2--chr2--name2")
)

chr_data_1 <- list(
  chromosomes = list("group1--1--chr1--name1", "group2--3--chr3--name3"),
  chromosome_groups = NA_character_
)
process_chromosomes(chr_data_1, select_input_data)

chr_data_2 <- list(
  chromosomes = NA,
  chromosome_groups = "group1"
)

process_chromosomes(chr_data_2, select_input_data)
```

`process_delimited_column`*Process Delimited Columns in Metadata*

Description

The specified column name is searched for across all the data frames. If the column is `pubmed_ids`, values are converted to integers.

Usage

```
process_delimited_column(  
  metadata,  
  column_name,  
  separator,  
  converters = DELIM_CONVERTERS  
)
```

Arguments

<code>metadata</code>	List. A list of data frames representing metadata sheets.
<code>column_name</code>	Character. The name of the column to process.
<code>separator</code>	Character. The delimiter used to split column values.
<code>converters</code>	Named list of functions. Specifies how to coerce columns with delimited strings into specific types or values. Defaults to <code>DELIM_CONVERTERS</code>

Value

A list of updated metadata with the specified column split into lists based on the delimiter and trimmed.

Examples

```
metadata <- list(  
  sheet1 = data.frame(pubmed_ids = c("123; 456", "130; 789; 102", NA))  
)  
  
process_delimited_column(metadata, "pubmed_ids", ";")
```

rollback_submission *Rollback Submission Endpoints and Log Responses*

Description

Rolls back specified endpoints for a submission identified by its accession ID using the client and logs the responses if a logfile is specified.

Usage

```
rollback_submission(id, endpoints, client = NULL, logfile = NULL, ...)
```

Arguments

id	A string representing the submission identifier. Must be an accession ID.
endpoints	A character vector of endpoint names to rollback.
client	List of functions. EGA API client created by <code>create_client</code> function from EGA API schema with put methods and rollback operations. If NULL, default client will be created by <code>create_client(extract_api())</code> . Defaults to NULL.
logfile	A string specifying the path to a log file. If NULL, no log is written. Defaults to NULL.
...	Additional arguments for future extensions (currently unused).

Value

A list of responses from the rollback operations for each endpoint.

Examples

```
mock_client <- list(
  "put__submissions__accession_id__datasets__rollback" =
    function(id) list(status = "rolled back")
)
rollback_submission("EGAB0000000001", list("datasets"), mock_client)
```

row_table_formatter *Format a Row Table*

Description

Format a Row Table

Usage

```
row_table_formatter(tab, params)
```

Arguments

tab	Data frame. The input table from a submission metadata file
params	List. Additional parameters for formatting. Takes a formatter params value from parser parameter yaml file.

Value

A cleaned and formatted tibble with correctly organized rows and columns, whitespace trimmed, and folding applied to specified columns.

Examples

```
# Formatter parameters
params <- list(fold = "extra_attributes")

# Sample data frame
df <- data.frame(
  ...1 = c("* Study", "* Title", "Extra Attributes", "Extra Attributes"),
  ...2 = c("Study1", "Title1", "A", "B"),
  ...3 = c("* Study", "* Title", "Extra Attributes", NA),
  ...4 = c("Study2", "Title2", "C", NA)
)

row_table_formatter(df, params)
```

samples_in_db

Check if sample aliases exist in the EGA database

Description

Validates uniqueness of sample aliases by comparing input against existing records in the EGA database. Throws an error if duplicates are found and retrieval is not enabled.

Usage

```
samples_in_db(samples, client = NULL, retrieve = FALSE)
```

Arguments

samples	Character vector of sample aliases to check.
client	An EGA API client object. If NULL, one is created.
retrieve	Logical scalar. If TRUE, exists without error even if samples are found in the database.

Value

Logical TRUE if validation passes.

Examples

```
my_client <- list(
  get__samples = function(prefix = NULL) {
    data.frame(alias = c("unique_sample1", "unique_sample_2"))
  }
)

samples_in_db(c("sample1", "sample2"), client = my_client, retrieve = FALSE)
```

save_log	<i>Save API Responses to a Log File</i>
----------	---

Description

This function saves a list of API responses to a specified log file in YAML format.

Usage

```
save_log(responses, logfile)
```

Arguments

responses	A list of responses to be saved.
logfile	A string specifying the path to the log file. If NULL, no file is written.

Value

Invisibly returns NULL

Examples

```
responses <- list(status = "success", data = list(a = 1, b = "text"))
save_log(responses, logfile = NULL)
```

step_msg	<i>Generate a Step-by-Step Message Function</i>
----------	---

Description

Creates a closure function to display sequential progress messages for a specified number of steps.

Usage

```
step_msg(steps)
```

Arguments

steps An integer specifying the total number of steps.

Value

A function that takes a message string as input and displays it along with the current step and total steps. The step count increments automatically with each call.

Examples

```
stepper <- step_msg(3)
stepper("Initializing") # "Step 1/3 - Initializing"
stepper("Processing") # "Step 2/3 - Processing"
stepper("Finalizing") # "Step 3/3 - Finalizing"
```

submit_table	<i>Submit a Data Frame to an API Endpoint Row by Row</i>
--------------	--

Description

This function iterates over rows of a data frame, submitting each row to a specified API endpoint function, and combines the responses into a single data structure.

Usage

```
submit_table(tab, id, endpoint_func)
```

Arguments

tab A data frame containing the data to be submitted.
id An EGA accession/provisional ID passed to the endpoint_func.
endpoint_func A function that handles the API request. It should accept id and a JSON body as arguments.

Value

Data frame. A combined response object from the API.

Examples

```
tab <- data.frame(a = 1:2, b = c("x", "y"))
mock_endpoint <- function(id, body) list(id = id, body = body)
submit_table(tab, 12345, mock_endpoint)
```

try_step

Execute a submission step with error handling and rollback

Description

Wraps a logic function in a tryCatch block to handle errors during a specific submission step, optionally triggering a rollback function.

Usage

```
try_step(step_name, logic_fn, rollback_fn, responses, logfile)
```

Arguments

step_name	Character. The name of the current workflow step.
logic_fn	Function. The primary logic to execute for this step.
rollback_fn	Function. A function to clean up if an error occurs.
responses	List. Current collection of API responses for logging.
logfile	Character. Path to the log file.

Value

The result of logic_fn().

Examples

```
try_step(
  "test", function() 1 + 1, function() print("fail"), list(), "log.txt"
)
```

`unbox_list`*Convert a List to an Unboxed JSON-Compatible Data Frame*

Description

Converts a list into a single-row data frame with unboxed elements if all elements have a length of 1. Otherwise, an error is raised.

Usage

```
unbox_list(l)
```

Arguments

`l` A list where all elements must have a length of 1.

Value

A data frame with unboxed elements, suitable for JSON conversion.

Examples

```
input_list <- list(a = 1, b = "text", c = TRUE)
unbox_list(input_list)
```

`unbox_row`*Convert a Data Frame Row to an Unboxed JSON Object*

Description

This function converts a single row of a data frame into an unboxed JSON object, effectively removing the array structure.

Usage

```
unbox_row(row)
```

Arguments

`row` A single row of a data frame.

Value

A JSON object with unboxed values for the input row.

Examples

```
row <- data.frame(a = 1, b = "text", stringsAsFactors = FALSE)[1, ]
unbox_row(row)
```

use_submission	<i>Retrieve or Delete Submission Data</i>
----------------	---

Description

Handles retrieval or deletion of data associated with a submission accession/provisional ID using a specified client and method.

Usage

```
use_submission(id, method, client = NULL)
```

Arguments

id	Character or numeric. Represents the submission identifier. Can be either an accession or provisional ID.
method	A string specifying the operation to perform. Valid options are "get" or "delete".
client	List of functions. EGA API client created by create_client function from EGA API schema with get and delete methods. If NULL, default client will be created by create_client(extract_api()). Defaults to NULL.

Value

A named list containing responses for datasets, analyses, runs, experiments, samples, and studies.

Examples

```
mock_client <- list(
  "get__submissions__accession_id__datasets" = function(id) {
    list(data = id)
  },
  "delete__submissions__provisional_id__datasets" =
    function(id) list(status = "deleted")
)
use_submission("EGAB12345678901", "get", mock_client)
```

validate_schema	<i>Validate a Payload Against a JSON Schema</i>
-----------------	---

Description

Function handles oneOf directives in a way that in a case of validation fail, it displays the overall result of the validation as first and then it tests separately against all oneOf sub schemas.

Usage

```
validate_schema(payload, schema)
```

Arguments

payload	The payload to validate against the schema. JSON string or single row of data frame converted to JSON representation with <code>unbox_row</code> function or a list with all items of length 1 converted to JSON representation with <code>unbox_list</code> function.
schema	List. The JSON schema defining the validation rules.

Value

Logical value indicating whether the payload is valid. If invalid, the result includes an errors attribute detailing the validation errors.

Examples

```
schema <- list(
  type = "object",
  properties = list(
    id = list(type = "integer"),
    title = list(type = "string")
  ),
  required = c("id")
)

payload_true <- data.frame(id = c(12345), title = c("abcd"))
payload_false <- data.frame(id = c("12345"), title = c(0.355))

validate_schema(jsonlite::unbox(payload_true), schema)
validate_schema(jsonlite::unbox(payload_false), schema)
```

validation_to_msg *Convert Validation Results to a Message*

Description

Convert Validation Results to a Message

Usage

```
validation_to_msg(v)
```

Arguments

v Logical. The validation result, which may include an `errors` attribute detailing validation errors.

Value

A character string summarizing the validation results. If validation errors are present, they are included in the message; otherwise, a success message is returned.

Examples

```
validation_result <- FALSE
attr(validation_result, "errors") <- data.frame(
  field = c("name"),
  message = c("Missing")
)
msg <- validation_to_msg(validation_result)
message(msg)
```

workflow_error_handler *Workflow Error Handler*

Description

Creates a custom error handler for managing errors during a workflow step. Logs responses, executes additional expressions, and stops execution with a detailed message and a stack trace.

Usage

```
workflow_error_handler(step, responses, logfile, ...)
```

Arguments

<code>step</code>	A string representing the current workflow step.
<code>responses</code>	A list of responses to be logged in case of an error.
<code>logfile</code>	A string specifying the path to the log file. If NULL, no file is written.
<code>...</code>	Additional expressions to evaluate when an error occurs.

Value

A function to handle errors during the specified workflow step.

Examples

```
handler <- workflow_error_handler(  
  step = "submission",  
  responses = list(),  
  logfile = NULL  
)  
  
tryCatch("Example code without error", error = handler)
```

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