Package ‘sbtools’

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Title USGS ScienceBase Tools

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Imports jsonlite, curl, httr (>= 1.0.0), methods, mime, utils, cli, keyring, tools

Suggests testthat, xml2, sf, knitr, rmarkdown

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URL https://github.com/DOI-USGS/sbtools,
https://doi-usgs.github.io/sbtools/

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Description

This package provides a rich interface to USGS’s ScienceBase [https://www.sciencebase.gov/](https://www.sciencebase.gov/) - a data cataloging and collaborative data management platform. For further information, see the sbtools manuscript [here](https://github.com/DOI-USGS/sbtools/issues).

Functions are included for searching for data, retrieving, creating, and updating datasets.

Details

Functionality in this package allows all users to query ScienceBase for data using a variety of metadata types (query_sb_text, query_sb_doi, query_sb_spatial). Items and associated information can be requested by item_get including item parents item_get_parent and children item_list_children. Data and attached files can be accessed for all available items through provided functionality (e.g., item_file_download).

Authentication

See the function authenticate_sb to authenticate. You’ll be required to pass in your ScienceBase username and password.

Authenticated users can create, update, and remove items (item_list_children, item_list_children, item_create, item_update, item_rm).

Feedback

Report any feedback or bugs at [https://github.com/DOI-USGS/sbtools/issues](https://github.com/DOI-USGS/sbtools/issues)

authenticate_sb

Authenticate to SB for subsequent calls

Description

This connects to SB, authenticates and gets a session token for communicating with SB. If you do not supply a username or password, you will be prompted to enter them.

Usage

authenticate_sb(username, password)
folder_create

Arguments

username  Sciencebase username
password  Sciencebase password, prompts user if not supplied and no password is returned
by 'keyring::key_get("sciencebase", username)'. See keyring-package documentation for more details.

current_session  Return current cached session

Description

Returns the currently cached SB session token. If there is no authenticated session, returns NULL.
Emits a warning if the session has expired.

Usage

current_session()

Examples

session = current_session()
#null unless currently authenticated
session

folder_create  Create a folder

Description

Create a special kind of item on ScienceBase that is intended to be a "folder" that contains one
or more child items. This is similar to a standard item (item_create) but defaults to showing
child-items on the ScienceBase web interface.

Usage

folder_create(parent_id = user_id(), name, ...)

Arguments

parent_id  An sbitem object or character ScienceBase ID corresponding to the parent item
(folder)
name  (character) the folder name
...  Additional parameters are passed on to GET, POST, HEAD, PUT
**identifier_exists**

**Value**

A response object

**Examples**

```r
## Not run:
folder_create(name="foobar345")

## End(Not run)
```

---

**identifier_exists**  
**Check if identifier exists**

**Description**

This function quickly checks to see if an identifier exists. It does a quick head request to skip the overhead of item metadata retrieval. This will also return FALSE if the identifier exists but is associated with an item that is unavailable due to permission restrictions.

**Usage**

```r
identifier_exists(sb_id, ...)
```

**Arguments**

- `sb_id` An sbitem object or a character ScienceBase ID corresponding to the item
- `...` Additional parameters are passed on to `GET`, `POST`, `HEAD`, `PUT`.

**Value**

Logical, TRUE or FALSE

**Examples**

```r
# identifier exists
identifier_exists(sb_id = "57976a0ce4b021cadec97890")

# identifier does not exist
identifier_exists(sb_id = "aaaaaaaaakkkkkkkbbbbbb")
```
initialize_sciencebase_session

Initialize ScienceBase Session

Description

Unless ‘token_text’ is provided, will open a browser for two factor authentication. Once logged in, retrieve the token from the user drop down in the upper right hand corner of the browser. Click the icon with the silhouette of a person, and select ‘Copy API Token.’ The token should be pasted into the popup prompt.

Usage

initialize_sciencebase_session(username = NULL, token_text = NULL)

Arguments

username email address of sciencebase user. Will be retrieved from the ‘sb_user’ environment variable if set. A prompt will be raised if not provided.

token_text character json formatted token text. ‘token_text’ is stashed in R_user_dir and does not need to be re-entered unless it becomes stale. If the token text is provided as input, no popup prompt will be raised.

is_logged_in

Check whether you’re logged into a ScienceBase session

Description

Check whether you’re logged into a ScienceBase session

Usage

is_logged_in()

Value

Logical, TRUE or FALSE

Examples

## Not run:
is_logged_in()

## End(Not run)
Description

A method to create multiple ScienceBase items with a single call and a single HTTP service request. Can be useful for improving performance of creating a large number of items at once.

Usage

```r
items_create(parent_id = user_id(), title, ..., info = NULL)
```

Arguments

- **parent_id**: An `sbitem` object or character ScienceBase ID corresponding to the parent item (folder). This must be of length 1 or more. If length 1, then we recycle it for every item.
- **title**: Two or more titles for the new SB items
- ..., **info**: (optional) list of metadata info for the new items. for each item include a named list of variables

Details

The length of the `title` and `info` values must be the same length - however, the `parent_id` can be of length 1 or equal to the length of each of `title` and `info` parameters

Value

One or more objects of class `sbitem` in a list

Examples

```r
## Not run:
# helper function to make a random name
aname <- function() paste0(sample(letters, size = 5, replace = TRUE), collapse = "")

# Create some items - by default we use your user ID
items_create(title = c(aname(), aname()))

# add additional items in the info parameter - by default we use your user ID
items_create(title = c(aname(), aname()),
             info = list(
                list(contacts = list(list(name = "Suzy")),
                list(contacts = list(list(name = "Brandy")))))
             )
```
# another example with more information - by default we use your user ID
items_create(title = c(aname(), aname()),
info = list(
  list(contacts = list(list(name = "Suzy"))),
  list(contacts = list(list(name = "Brandy")))
))

# Pass an object of class sbitem
(x <- folder_create(user_id(), aname()))
items_create(x, title = c(aname(), aname()))

## End(Not run)

## items_update
Update many SB items with new metadata

Description

A method to update multiple ScienceBase items with a single call and a single HTTP service request. Can be useful for improving performance of updating a large number of items at once.

Usage

items_update(sb_id, info, ...)

Arguments

- sb_id: An sbitem object or a character ScienceBase ID corresponding to the item
- info: list of metadata info (key-value pairs) to change on the item
- ...: Additional parameters are passed on to PUT

Details

If length of sb_id > 1, then length of info input must be the same

Value

One or more objects of class sbitem in a list

Examples

## Not run:
# helper function to make a random name
aname <- function() paste0(sample(letters, size = 5, replace = TRUE), collapse = "")

res <- items_create(user_id(), title = c(aname(), aname()))
out <- items_update(res, info = list( list(title = aname()), list(title = aname()) ) )
vapply(out, "[[", ",", "title")
items_upsert  

**Upsert many SB items**

**Description**

Either creates or updates (if items already exist)

**Usage**

```r
items_upsert(parent_id = user_id(), title = NULL, ..., info = NULL)
```

**Arguments**

- `parent_id`  
  An `sbitem` object or character ScienceBase ID corresponding to the parent item (folder)
- `title`  
  The title of the new SB item
- `...`  
  Additional parameters are passed on to `GET`, `POST`, `HEAD`, `PUT`
- `info`  
  (optional) list of metadata info for the new item

**Value**

An object of class `sbitem`

**Examples**

```r
## Not run:  
# helper function to make a random name  
aname <- function() paste0(sample(letters, size = 5, replace = TRUE), collapse = "")

# Create some item - by default we use your user ID  
z1 <- item_create(title = aname())  
z2 <- item_create(title = aname())

# Upsert items  
(x <- items_upsert(list(z1, z2), title = c(aname(), aname())))

# Call item_upsert again, updates this time  
items_upsert(x, info = list(  
  contacts = list(list(name = "Suzy"))
))
```

## End(Not run)
item_append_files  
*Upload File to Item*

**Description**

Adds a file to an item

**Usage**

```r
item_append_files(sb_id, files, ..., scrape_files = TRUE)
```

**Arguments**

- `sb_id` An `sbitem` object or a character ScienceBase ID corresponding to the item
- `files` A string vector of paths to files to be uploaded
- `...` Additional parameters are passed on to `GET`, `POST`, `HEAD`, `PUT`.
- `scrape_files` logical should the files be scraped for metadata? If TRUE, sciencebase will attempt to create extensions based on the files. For example, for shapefiles, this will result in a shapefile extension to be returned as a facet of the sciencebase item. See item: "58069258e4b0824b2d1d422e" for an example.

**Value**

An object of class `sbitem`

**Examples**

```r
## Not run:
res <- item_create(user_id(), "testing 123")
cat("foo bar", file = "foobartxt")
item_append_files(res$id, "foobartxt")
item_rm(res)

## End(Not run)
```

---

**item_create**  
*Create a new SB item*

**Description**

Create a new item on ScienceBase with the requested parent and item title. Info can be provided to populate metadata at the time of creation.
item_create

Usage

item_create(parent_id = user_id(), title, ..., info)

Arguments

parent_id  An sbitem object or character ScienceBase ID corresponding to the parent item (folder)
title  The title of the new SB item...
info  (optional) list of metadata info for the new item

Value

An object of class sbitem

Examples

## Not run:
# Create an item - by default we use your user ID
item_create(title = "testing 123")

# Pass an object of class sbitem
x <- folder_create(user_id(), "foobar456")
item_create(x, "foobar456-item")

## End(Not run)

item_exists

check if identifier tuple already exists on SB

Description

returns TRUE if tuple already belongs to a sciencebase item, FALSE if not

Usage

item_exists(scheme, type, key, ...)

Arguments

scheme  the identifier scheme
type  the identifier type
key  the identifier key...

...  Additional parameters are passed on to GET
item_file_download

Value

  boolean for whether item exists

Examples

## Not run:
item_exists('mda_streams','ts_doobs','nwis_01018035')
item_exists('mda_streams','site_root','nwis_01018035')

## End(Not run)

---

item_file_download  Download files attached to item

Description

Function to download files attached to an item on SB. Either files can be specified directly using the names and destinations parameters, or a dest_dir can be supplied where all attached files will be written with the names as stored on SB.

Usage

item_file_download(
  sb_id,
  ...,
  names,
  destinations,
  dest_dir = getwd(),
  overwrite_file = FALSE
)

Arguments

  sb_id  An sbitem object or a character ScienceBase ID corresponding to the item
  ...    Additional parameters are passed on to GET, POST, HEAD, PUT.
  names  String vector list of file names attached to item that you wish to download.
  destinations  String vector list of destinations for requested files. Must be same length as names
  dest_dir  A directory path for saving files when names destinations parameter is not specified.
  overwrite_file  Boolean indicating if file should be overwritten if it already exists locally

Value

  Character vector of full paths to local files
Examples

## Not run:

```r
#downloads all files attached to this item
item_file_download('627f1572d34e3bef0c9a30d8', dest_dir=tempdir())

#downloads a specific file attached to this item
item_file_download('627f1572d34e3bef0c9a30d8', names='example.txt',
                  destinations=file.path(tempdir(), 'out.txt'))
```

## End(Not run)

---

**item_get**

*Retrieve SB item*

Description

Retrieves an item and its metadata from ScienceBase based on its unique ID. Errors if the requested item ID does not exist or access is restricted due to permissions.

Usage

```r
item_get(sb_id, ...)
```

Arguments

- `sb_id`:
  - An `sbitem` object or a character ScienceBase ID corresponding to the item
- `...`:
  - Additional parameters are passed on to `GET, POST, HEAD, PUT`.

Value

An object of class `sbitem`

Examples

```r
# Get an item
item_get("4f4e4b24e4b07f02db6ae14")

# Search for item IDs, then pass to item_get
library("httr")
res <- query_items(list(s = "Search", q = "water", format = "json"))
if(inherits(res, "response") && res$status != 404) {
  ids <- vapply(httr::content(res)$items, "+\[", "," , "id\"
                  lapply(ids[1:3], item_get)
}
item_get_fields  Retrieve specific fields from an SB item

Description

Retrieve specific fields from an SB item

Usage

item_get_fields(sb_id, fields, ..., drop = TRUE)

Arguments

- **sb_id**: An sbitem object or a character ScienceBase ID corresponding to the item
- **fields**: a vector of fields
- **...**: Additional parameters are passed on to GET, POST, HEAD, PUT.
- **drop**: logical. If only one field is selected, should the list format be dropped?

Value

List serialization of chosen metadata for an SB item

Examples

# Get certain fields from an item
item_get_fields("63cb38b2d34e06fef14f40ad", c('title', 'citation', 'contacts'))

# If only 1 field selection, do or don't drop list format
item_get_fields("63cb38b2d34e06fef14f40ad", 'title')
item_get_fields("63cb38b2d34e06fef14f40ad", 'title', drop = FALSE)

item_get_parent  Get an item's parent ID

Description

Retrieves the parent of a supplied item based on the ScienceBase item tree hierarchy.

Usage

item_get_parent(sb_id, ...)
**item_list_children**

**Arguments**

- **sb_id**
  - An `sbitem` object or a character ScienceBase ID corresponding to the item
- ...
  - Additional parameters are passed on to `GET`, `POST`, `HEAD`, `PUT`.

**Value**

- An item object representing the parent of the supplied item.

**Examples**

```r
item_get_parent("57976a0ce4b021cadec97890")
item_get_parent(item_get("57976a0ce4b021cadec97890"))
```

---

**Description**

Returns a list of child IDs for a ScienceBase item

**Usage**

```r
item_list_children(sb_id, fields = c("id", "title"), ..., limit = 20)
```

**Arguments**

- **sb_id**
  - An `sbitem` object or a character ScienceBase ID corresponding to the item
- **fields**
  - A character vector of requested data fields. Defaults to ‘id’ and ‘title’. Full list of possible fields is available online in SB documentation.
- ...
  - Additional parameters are passed on to `GET`, `POST`, `HEAD`, `PUT`.
- **limit**
  - Max children returned.

**Value**

- List of `sbitem` for each child item.
Examples

## Not run:
item_list_children(user_id())
## End(Not run)

item_list_children(as.sbitem('5060b03ae4b00fc20c4f3c8b'))
item_list_children(item_get('5060b03ae4b00fc20c4f3c8b'))

---

**item_list_files**  
*Get list of files attached to SB item*

Description

Lists all files attached to a SB item. Files can be downloaded from ScienceBase using `item_file_download`.  
(advanced) Recursive options lists all files attached to an item and all children items.  
NOTE: A sciencebase item can contain so-called "extensions". The sciencebase item data model refers to the information that describes an extension as a "facet". Some extension facets contain files (such as with a shapefile). The "facet" attribute of the return from this function will contain the name of the facet the file came from if the file was found in a facet.

Usage

```r
item_list_files(sb_id, recursive = FALSE, fetch_cloud_urls = TRUE, ...)
```

Arguments

- `sb_id`: An `sbitem` object or a character ScienceBase ID corresponding to the item  
- `recursive` (logical): List files recursively. Default: `FALSE`  
- `fetch_cloud_urls` (logical): fetch a tokenized cloud download URLs? Default: `TRUE` This option will take slightly longer but the ‘url’ attribute of the returned list will work for direct file downloads or use with pther applications and libraries.  
- `...`: Additional parameters are passed on to `GET, POST, HEAD, PUT`.

Value

A data.frame with columns `fname`, `size`, `url`, and `facet`. If item has no attached files, returns a zero row data.frame.
## Examples

```r
## Not run:

# regular files
item_list_files("57976a0ce4b021cadec97890")

# files in facets
item_list_files("Sebe92af82ce476925e44b8f")

# list files recursively
## create item
id <- item_create(user_id(), title="some title")
## 1. create nested item w/ file
file <- system.file("examples", "books.json", package = "sbtools")
id2 <- item_create(id, title = "newest-thing")
item_upload_create(id2, file)
## 2. create nested item w/ file
file <- system.file("examples", "species.json", package = "sbtools")
id3 <- item_create(id, title = "a-new-thing")
item_upload_create(id3, file)
## 3. create nested item w/ file
file <- system.file("examples", "data.csv", package = "sbtools")
id4 <- item_create(id, title = "another-thing")
item_upload_create(id4, file)
item_list_files(id = '56562348e4b071e7ea53e09d', recursive = FALSE) # default
item_list_files(id = '56562348e4b071e7ea53e09d', recursive = TRUE)

## End(Not run)
```

---

**item_move**

*Move item from one folder to another*

**Description**

Move item from one folder to another

**Usage**

```r
item_move(sb_id, id_new, ...)
```

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>sb_id</td>
<td>An sbitem object or a character ScienceBase ID corresponding to the item</td>
</tr>
<tr>
<td>id_new</td>
<td>Folder/item to move id to. A ScienceBase ID or something that can be coerced to a SB item ID by as.sbitem</td>
</tr>
<tr>
<td>...</td>
<td>Additional parameters are passed on to GET, POST, HEAD, PUT.</td>
</tr>
</tbody>
</table>
item_publish_cloud

Value

An object of class sbitem. Same as id, but with new parent id

Examples

```r
## Not run:
# create 1st folder
(fold1 <- folder_create(user_id(), "bear123"))
(res <- item_create(fold1, "item-to-move"))

# create 2nd folder
(fold2 <- folder_create(user_id(), "bear456"))

# move item in 1st folder to 2nd folder
(res2 <- item_move(res, fold2))

# test identical
identical(res2$parentId, fold2$id)

## End(Not run)
```

---

**item_publish_cloud**   *Publish file to public cloud S3 bucket*

Description

moves a cloud file from the S3 bucket only available via ScienceBase authenticated services to a public S3 bucket.

Usage

`item_publish_cloud(sb_id, files, ...)`

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>sb_id</td>
<td>An sbitem object or a character ScienceBase ID corresponding to the item</td>
</tr>
<tr>
<td>files</td>
<td>A string vector of paths to files to be uploaded</td>
</tr>
<tr>
<td>...</td>
<td>Additional parameters are passed on to <code>GET, POST, HEAD, PUT</code>.</td>
</tr>
</tbody>
</table>

Value

web service response invisibly.
item_rename_files

Examples

```r
## Not run:
res <- item_create(user_id(), "testing 123")
cat("foo bar", file = "foobar.txt")
item_upload_cloud(res$id, "foobar.txt")
item_publish_cloud(res$id, "foobar.txt")

## End(Not run)
```

---

**item_rename_files**  Rename item attached files

**Description**

Renames files attached to an SB item.

**Usage**

```r
item_rename_files(sb_id, names, new_names, ...)
```

**Arguments**

- `sb_id` An sbitem object or a character ScienceBase ID corresponding to the item
- `names` List of names of files to rename
- `new_names` List of new file names to use
- `...` Additional parameters are passed on to `GET`, `POST`, `HEAD`, `PUT`.

**Examples**

```r
## Not run:
names = c('file1.txt', 'file2.txt')
new_names = c('newname1.txt', 'newname2.txt')
item_rename_files('sbid', names, new_names)

## End(Not run)
```
**item_replace_files**  
*Replace files associated with an item*

**Description**

replaces existing files associated with an item with a new one.

NOTE: This function will not replace files stored in facets. Until and if facet support is added, direct alteration of the science base item object is required to manipulate facets.

**Usage**

```r
item_replace_files(sb_id, files, ..., all = FALSE, scrape_files = FALSE)
```

**Arguments**

- `sb_id`: An `sbitem` object or a character ScienceBase ID corresponding to the item
- `files`: A character vector of file paths
- `...`: Additional parameters are passed on to `GET`, `POST`, `HEAD`, `PUT`.
- `all`: A boolean indicating if all attached files should be removed before uploading new files. FALSE if only files with matching names should be replaced. If you wish to upload files with duplicate names, see `item_append_files`. Defaults to FALSE.
- `scrape_files`: logical should the files be scraped for metadata? If TRUE, sciencebase will attempt to create extensions based on the files. For example, for shapefiles, this will result in a shapefile extension to be returned as a facet of the sciencebase item. See item: "58069258e4b0824b2d1d422e" for an example.

**item_rm**  
*Remove item from SB*

**Description**

Remove an item from ScienceBase. This is not reversible and will delete an item and its attached files. (advanced) Recursive is to be used with care and could result in unexpected file deletion.

**Usage**

```r
item_rm(sb_id, ..., limit = 1000, recursive = FALSE)
```
**Argument**

- **sb_id**
  - An `sbitem` object or a character ScienceBase ID corresponding to the item
  - Additional parameters are passed on to `GET`, `POST`, `HEAD`, `PUT`.
- **limit**
  - The maximum number of child items to remove when called with `recursive=TRUE`.
- **recursive**
  - Logical, `FALSE` by default. **CAUTION:** Setting `recursive=TRUE` means that not only will this item be deleted, but so will all its child items and their child items and so on.

**Value**

An `httr` response object

**Examples**

```r
## Not run:
res <- item_create(user_id(), "item-to-delete")
item_rm(res)
## End(Not run)
```

---

**Description**

Removes existing files associated with an item.

**NOTE:** This function will not alter facets which can also contain facets. To manipulate facets, the facet element of a sciencebase item must be altered and updated with `item_update`.

This function is the key way to remove files attached to SB items.

**Usage**

```
item_rm_files(sb_id, files, ...)
```

**Arguments**

- **sb_id**
  - An `sbitem` object or a character ScienceBase ID corresponding to the item
- **files**
  - A character vector of file names to remove. If not supplied, defaults to removing all attached files.
  - Additional parameters are passed on to `GET`, `POST`, `HEAD`, `PUT`.

**Value**

An updated object of class `sbitem`
Examples

```r
## Not run:
res <- item_create(user_id(), "item456")
cat("foo bar", file = "foobar.txt")
item_append_files(res, "foobar.txt")
res <- item_get(res)
res$files[[1]]$name
res2 <- item_rm_files(res)
res2$files

## End(Not run)
```

---

**item_update**

Update a SB item with new metadata

**Description**

Updates metadata associated with a ScienceBase item based on supplied list of new or updated metadata elements.

**Usage**

```r
item_update(sb_id, info, ...)
```

**Arguments**

- `sb_id` An `sbitem` object or a character ScienceBase ID corresponding to the item
- `info` list of metadata info (key-value pairs) to change on the item
- `...` Additional parameters are passed on to `GET`, `POST`, `HEAD`, `PUT`.

**Value**

An object of class `sbitem`

**Examples**

```r
## Not run:
res <- item_create(user_id(), "item-to-update")
out <- item_update(res, list(title = "item-updated"))
out$title

## End(Not run)
```
**item_update_identifier**

*Add custom identifier to an existing item*

**Description**

Adds or updates an item’s alternative identifier. This can add additional identifiers or update those already in place. See `query_item_identifier` for finding items based on alternative identifier.

**Usage**

```r
item_update_identifier(sb_id, scheme, type, key, ...)
```

**Arguments**

- `sb_id` An `sbitem` object or a character ScienceBase ID corresponding to the item
- `scheme` The identifier scheme
- `type` The identifier type
- `key` The identifier key
- `...` Additional parameters are passed on to `GET, POST, HEAD, PUT`.

**Examples**

```r
## Not run:
initialize_sciencebase_session()
item_update_identifier("5485fd99e4b02acb4f0c7e81", "scheme", "type", "key")

## End(Not run)
```

---

**item_upload_cloud** *Upload File to Item Cloud Storage*

**Description**

Adds a file to an item in cloud storage

**Usage**

```r
item_upload_cloud(sb_id, files, ..., status = TRUE)
```
Arguments

- **sb_id**: An `sbitem` object or a character ScienceBase ID corresponding to the item.
- **files**: A string vector of paths to files to be uploaded.
- **...**: Additional parameters are passed on to `GET`, `POST`, `HEAD`, `PUT`.
- **status**: Logical display upload status?

Value

Success message invisibly. NOTE: cloud processing can take some time so the added file may not appear immediately. For this reason, a sciencebase item json is NOT returned as is done with other similar functions.

Examples

```r
## Not run:
res <- item_create(user_id(), "testing 123")
cat("foo bar", file = "foobar.txt")
item_upload_cloud(res$id, "foobar.txt")
## End(Not run)
```

---

**Description**

Create a new item with files attached, all in one call to SB.

**Usage**

```r
item_upload_create(parent_id, files, ..., scrape_files = TRUE)
```

**Arguments**

- **parent_id**: An `sbitem` object or character ScienceBase ID corresponding to the parent item (folder).
- **files**: A string vector of paths to files to be uploaded.
- **...**: Additional parameters are passed on to `GET`, `POST`, `HEAD`, `PUT`.
- **scrape_files**: Logical should the files be scraped for metadata? If TRUE, sciencebase will attempt to create extensions based on the files. For example, for shapefiles, this will result in a shapefile extension to be returned as a facet of the sciencebase item. See item: "58069258e4b0824b2d1d422e" for an example.
**item_upsert**

| **Upsert an SB item** |

**Description**

Either creates or updates (if item already exists)

**Usage**

`item_upsert(parent_id = user_id(), title = NULL, ..., info = NULL)`

**Arguments**

- `parent_id`: An `sbitem` object or character ScienceBase ID corresponding to the parent item (folder)
- `title`: The title of the new SB item
- `...`: Additional parameters are passed on to `GET, POST, HEAD, PUT`
- `info`: (optional) list of metadata info for the new item

**Value**

An object of class `sbitem`

**Examples**

```r
## Not run:
# helper function to make a random name
aname <- function() paste0(sample(letters, size = 5, replace = TRUE), collapse = "")

# Create an item - by default we use your user ID
(x <- item_upsert(title = aname()))

# Call item_upsert again, updates this time
item_upsert(x, info = list(
    contacts = list(list(name = "Suzy")))
```
query_items

Description
Query SB for items using generic query parameters

Usage
query_items(query_list, ...)

Arguments
query_list List of item query selectors. See Details.
... Additional parameters are passed on to GET

Details
The following is a list of query parameters you can use in the query_list parameter.

- s (character): Only option: "Search"
- format (character): One of "json", "xml", "csv", or "atom"
- q (character): Query string
- q (character): Lucene query string
- max (integer): Number of records to return. Default: 20
- offset (integer): Record to start at. Default: 1
- fields (character): Character vector of fields to return
- folderId (character): Alphanumeric string representing folder ID
- parentId (character): Alphanumeric string representing folder ID. This can be used to return all children items within the folder, but not within sub-folders.
- sort (character) One of "firstContact", "dateCreated", "lastUpdated", or "title". By default sorted by search score
- order (character) One of "asc" or "desc"
- ids Vector of item ids.
- ancestors (character): Alphanumeric string representing folder ID. This can be used to return all children items within the folder, even within sub-folders. Used as a filter
- tags Filter by tags, e.g., "distribution". Used as a filter
- browseCategory One of .... Used as a filter
• browseType One of .... Used as a filter
• dateRange A json string with keys dateType and choice. Where dateType is one of Acquisition, Award, Collected, dateCreated, Received, Reported, Transmitted, Due, End, Info, lastUpdated, Publication, Release, or Start. And where choice is one of day, week, month, year, or range (if range selected, also supply start and end keys with dates of the form YYYY-MM-DD). Used as a filter
• projectStatus One of Active, Approved, Completed, In Progress, Proposed. Used as a filter
• spatialQuery A WKT string. Used as a filter
• extentQuery Use existing extents (footprints) to search against item bounding boxes and representational points. This is an alphanumeric string.

Value
An object of class response

See Also
query_item_identifier, query_item_in_folder

Examples
## Not run:
# Basic query
library("httr")
res <- query_items(list(s = "Search", q = "water", format = "json"))
httr::content(res)

# Paging
## max - number of results
res <- query_items(list(s = "Search", q = "water", format = "json", max = 2))
length(httr::content(res)$items)
res <- query_items(list(s = "Search", q = "water", format = "json", max = 30))
length(httr::content(res)$items)
## offset - start at certain record
res <- query_items(list(s = "Search", q = "water", format = "json",
 max = 30, offset = 10))
httr::content(res)
## links - use links given in output for subsequent queries
httr::content(httr::GET(
 content(res)$nextLink$url
 ))

# Return only certain fields
res <- query_items(list(s = "Search", q = "water", format = "json", fields = 'title'))
httr::content(res)$items[[1]]

# Search a folder ID
res <- query_items(list(s = "Search", q = "water", format = "json",
folderId = '504216b9e4b04b508bfd337d'))
httr::content(res)$items
# Filter by ancestor
query_items(list(s = "Search", ancestors = "4f831626e4b0e84f6086809b", format = "json"))

# Filter by tags
content(query_items(list(s = "Search", tags = "distribution", format = "json")))

# Filter by browse category
content(query_items(list(s = "Search", browseCategory = "Image", format = "json")))

# Filter by browse type
content(query_items(list(s = "Search", browseType = "Collection", format = "json")))

# Filter by WKT geometry string
wkt1 <- "POLYGON((-104.4 41.0,-95.1 41.0,-95.1 37.5,-104.4 37.5,-104.4 41.0))"
wkt2 <- "POLYGON((-104.4 38.3,-95.2 38.3,-95.2 33.7,-104.4 34.0,-104.4 38.3))"
content(query_items(list(s = "Search", spatialQuery = wkt1, format = "json")))
content(query_items(list(s = "Search", spatialQuery = wkt1, spatialQuery = wkt2, format = "json")))

# Project status
content(query_items(list(s = "Search", projectStatus = "Active", format = "json")))

# Date range
query_items(list(s = "Search", dateRange = '{"dateType":"Collected","choice":"year"}', format = "json"))
query_items(list(s = "Search", dateRange = '{"dateType":"lastUpdated","choice":"month"}', format = "json"))
query_items(list(s = "Search", dateRange = '{"dateType":"Release","choice":"range","start":"2014-09-01","end":"2015-09-01"}', format = "json"))

# Extent query
## just an alphanumeric code
content(query_items(list(s = "Search", extentQuery = '2873462', format = "json")))
## with buffering, intersect
content(query_items(list(s = "Search", extentQuery = '{"extent":2873462, "relation":"intersects","buffer":"5"}', format = "json")))
## with buffering, within
content(query_items(list(s = "Search", extentQuery = '{"extent":2873462, "relation":"within","buffer":"5"}', format = "json")))
## with buffering, within
content(query_items(list(s = "Search", extentQuery = '{"extent":2873462, "relation":"disjoint","buffer":"5"}', format = "json")))

# Lucene query
## note, you have to pass the q parameter if you pass the lq parameter
content(query_items(list(s = "Search", q = '', lq = "sage OR grouse")))

## End(Not run)
query_item_identifier  Query SB for items based on custom identifier

Description

Find all items under a scheme or also query by for a specific type and key

Usage

query_item_identifier(scheme, type = NULL, key = NULL, ..., limit = 20)

Arguments

- **scheme**: The identifier scheme
- **type**: (optional) The identifier type
- **key**: (optional) The identifier key
- **...**: Additional parameters are passed on to GET
- **limit**: Max number of matching items to return

Value

The SB item id for the matching item. NULL if no matching item found.

Examples

```r
## Not run:
authenticate_sb()

ex_item = item_create(title="identifier example")
item_update_identifier(ex_item, "project1", "dataset1", "key1")
ex2_item = item_create(title="identifier example 2")
item_update_identifier(ex2_item, "project1", "dataset1", "key2")

#query the specific item
query_item_identifier("project1", "dataset1", "key1")

#or get the collection of items based on the ID hierarchy
query_item_identifier("project1")

item_rm(ex_item)
item_rm(ex2_item)

## End(Not run)
```
query_item_in_folder  Search within an SB folder

Description
Search for text in the title, abstract, etc. within an SB folder and any subfolders.

Usage
query_item_in_folder(text, folder, ..., limit = 20)

Arguments
- text: text in the title, abstract, etc. of the desired item
- folder: an SB item ID for the folder to search in
- ... Additional parameters are passed on to GET
- limit: Max number of matching items to return

Value
A list of matching items as sbitem objects.

query_sb  Query SB for items using generic query parameters

Description
Generic SB query function to construct advanced queries.
The following is a list of query parameters you can use in the query_list parameter.

- q (character): Query string
- q (character): Lucene query string
- fields (character): Character vector of fields to return
- folderId (character): Alphanumeric string representing folder ID
- parentId (character): Alphanumeric string representing folder ID. This can be used to return all children items within the folder, but not within sub-folders.
- sort (character) One of "firstContact", "dateCreated", "lastUpdated", or "title". By default sorted by search score
- order (character) One of "asc" or "desc"
- ids Vector of item ids.
- ancestors (character): Alphanumeric string representing folder ID. This can be used to return all children items within the folder, even within sub-folders. Used as a filter
• tags Filter by tags, e.g, "distribution". Used as a filter
• browseCategory One of .... Used as a filter
• browseType One of .... Used as a filter
• dateRange A json string with keys dateType and choice. Where dateType is one of Acquisition, Award, Collected, dateCreated, Received, Reported, Transmitted, Due, End, Info, lastUpdated, Publication, Release, or Start. And where choice is one of day, week, month, year, or range (if range selected, also supply start and end keys with dates of the form YYYY-MM-DD). Used as a filter
• projectStatus One of Active, Approved, Completed, In Progress, Proposed. Used as a filter
• spatialQuery A WKT string. Used as a filter
• extentQuery Use existing extents (footprints) to search against item bounding boxes and representational points. This is a alphanumeric string.

Usage

query_sb(query_list, ..., limit = 20)

Arguments

query_list List of item query selectors. See Details.
... Additional parameters are passed on to GET
limit Maximum number of returned items. Will do paging to retrieve results when limit is over 1000. Use with caution, queries 10k results are slow.

Value

A list of sbitem objects

See Also

query_items

Examples

## Not run:
query_sb(list(q = "water"))

# Search by project status
query_sb(list(projectStatus = "Active"))

# Search a folder ID
query_sb(list(q = "water", folderId = '504216b9e4b04b508bfd337d'))

# Filter by ancestor
query_sb(list(ancestors = "4f831626e4b0e84f6086809b"))

# Filter by tags
query_sb(list(tags = "distribution"))
# Filter by browse category
query_sb(list(browseCategory = "Image"))

# Filter by browse type
query_sb(list(browseType = "Map Service"))

# Filter by WKT geometry string
wkt1 <- "POLYGON((-104.4 41.0,-95.1 41.0,-95.1 37.5,-104.4 37.5,-104.4 41.0))"
wkt2 <- "POLYGON((-104.4 38.3,-95.2 38.3,-95.2 33.7,-104.4 34.0,-104.4 38.3))"
query_sb(list(spatialQuery = wkt1))
query_sb(list(spatialQuery = wkt1, spatialQuery = wkt2))

# Date range
query_sb(list(dateRange = '{"dataType":"Collected","choice":"year"}'))
query_sb(list(dateRange = '{"dataType":"lastUpdated","choice":"month"}'))
query_sb(list(dateRange = '{"dataType":"Release","choice":"range","start":"2014-09-01","end":"2015-09-01"}'))

## End(Not run)

---

query_sb_datatype  Query SB for specific data type

**Description**

Queries ScienceBase for items with matching datatype.

**Usage**

query_sb_datatype(datatype, ..., limit = 20)

**Arguments**

- **datatype**  Character string indicating datatype. See `sb_datatypes` for full list of available datatypes.
- **...**  Additional parameters are passed on to `GET`
- **limit**  Maximum number of returned items. Will do paging to retrieve results when limit is over 1000. Use with caution, queries 10k results are slow.

**Value**

A list of `sbitem` objects. List of length 0 means no matches were found.
query_sb_date

Examples

#query for items with WFS Layer data
query_sb_datatype('Static Map Image')

#query for US Topo maps
query_sb_datatype('Map Service')

query_sb_date Query SB for items within a date range

Description

Queries ScienceBase for items with timestamps within a certain date/time range.

Usage

query_sb_date(
  start = as.POSIXct("1970-01-01"),
  end = Sys.time(),
  date_type = "lastUpdated",
  ...
  limit = 20
)

Arguments

start Start date as POSIXct object. Defaults to 1970-01-01
end End date as POSIXct object. Defaults to today.
date_type Which object timestamp to query against. Options are (case sensitive): 'Acquisition', 'Award', 'Collected', 'dateCreated', 'Received', 'Reported', 'Transmitted', 'Due', 'End', 'Info', 'lastUpdated', 'Publication', 'Release', 'Repository Created', 'Repository Updated', 'Start'.
... Additional parameters are passed on to GET
limit Maximum number of returned items. Will do paging to retrieve results when limit is over 1000. Use with caution, queries 10k results are slow.

Examples

## Not run:
# find items updated today
query_sb_date(Sys.time(), Sys.time())

# find items with publications from the 1970's
query_sb_date(as.POSIXct('1970-01-01'), as.POSIXct('1980-01-01'),
query_sb_doi

```
date_type='Publication', limit=1000)
```

## End(Not run)

---

query_sb_doi  

*Query SB for specific DOI (Digital Object Identifier)*

**Description**

Queries for ScienceBase items with a specific DOI identifier. In ScienceBase, these are stored as additional unique identifiers.

**Usage**

```
query_sb_doi(doi, ..., limit = 20)
```

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>doi</strong></td>
<td>DOI to search for as character</td>
</tr>
<tr>
<td><strong>...</strong></td>
<td>Additional parameters are passed on to GET</td>
</tr>
<tr>
<td><strong>limit</strong></td>
<td>Maximum number of returned items. Will do paging to retrieve results when limit is over 1000. Use with caution, queries 10k results are slow.</td>
</tr>
</tbody>
</table>

**Value**

A list of *sbitem* objects. List of length 0 means no matches were found.

**Examples**

```
# Two example DOI-specific queries
query_sb_doi('10.5066/F7M043G7')

query_sb_doi('10.5066/F7Z60M35')
```
**query_sb_spatial**  
*Query SB based on spatial extent*

**Description**

Queries ScienceBase based on a spatial bounding box. Accepts either an sp spatial data object (uses the spatial object’s bounding box) or long/lat coordinates defining the bounding box limits.

**Usage**

```r
query_sb_spatial(bbox, long, lat, bb_wkt, ..., limit = 20)
```

**Arguments**

- **bbox**: An sf spatial data object. The bounding box of the object is used for the query.
- **long**: A vector of longitude values that will define the boundaries of a bounding box. Min and Max of supplied longitudes are used. (alternate option to bbox).
- **lat**: A vector of latitude values that will define the boundaries of a bounding box. Min and Max of supplied latitude are used. (alternate option to bbox).
- **bb_wkt**: A character string using the Well Known Text (WKT) standard for defining spatial data. Must be a POLYGON WKT object.
- **...**: Additional parameters are passed on to GET
- **limit**: Maximum number of returned items. Will do paging to retrieve results when limit is over 1000. Use with caution, queries 10k results are slow.

**Examples**

```r
#specify the latitude and longitude points to define the bounding box range.
# This is simply bottom left and top right points
query_sb_spatial(long=c(-104.4, -95.1), lat=c(37.5, 41.0), limit=3)

#use a pre-formatted WKT polygon to grab data
query_sb_spatial(bb_wkt="POLYGON((-104.4 41.0,-95.1 41.0,-95.1 37.5,-104.4 37.5,-104.4 41.0))", limit=3)
```
query_sb_text  

*Query SB for items containing specific text*

**Description**

Queries for ScienceBase items that have matching text in the title or description

**Usage**

```r
query_sb_text(text, ..., limit = 20)
```

**Arguments**

- `text` Text string for search
- `...` Additional parameters are passed on to `GET`
- `limit` Maximum number of returned items. Will do paging to retrieve results when limit is over 1000. Use with caution, queries 10k results are slow.

**Value**

A list of `sbitem` objects. List of length 0 means no matches were found.

**Examples**

```r
#query for a person's name
query_sb_text('Luna Leopold')

#query for one of the old river gaging stations
query_sb_text('Lees Ferry')
```

---

**sbitem  

*ScienceBase item class*

**Description**

ScienceBase item class

**Usage**

```r
as.sbitem(x, ...)
```

```r
## Default S3 method:
as.sbitem(x, ...)

is.sbitem(x)
```
sb_datatypes

Arguments

  x  Input, variety of things, character, list, or sbitem class object
  ...
  Further args passed on to item_get, only in the method for character class inputs

Examples

  # Single item from item_get()
  item_get("57976a0ce4b021cadec97890")

  # Get many w/ e.g., an lapply() call
  library("httr")
  res <- query_items(list(s = "Search", q = "water", format = "json"))
  if(res$status == 200) {
    ids <- vapply(httr::content(res)$items, "[", "", "id")
    (out <- lapply(ids[1:3], item_get))
  }
  # create item class from only an item ID
  as.sbitem("5ebe92af82ce476925e44b8f")

  # sbitem gives back itself
  (x <- as.sbitem("5ebe92af82ce476925e44b8f"))
  as.sbitem(x)

sb_datatypes  Query SB for all available datatypes

Description

Queries ScienceBase for the list of all available datatypes. This can be coupled with query_sb_datatype to query based on the type of data

Usage

sb_datatypes(limit = 50)

Arguments

  limit  Maximum number of returned items. Will do paging to retrieve results when limit is over 1000. Use with caution, queries 10k results are slow.

Examples

  ## Not run:
  #return all datatypes (limit 50 by default)
  sb_datatypes()

  ## End(Not run)
sb_ping  

*Ping ScienceBase to see if it's available*

**Description**

Ping ScienceBase to see if it’s available.

**Usage**

```
sb_ping(...)```

**Arguments**

...   Additional parameters are passed on to GET

**Value**

Boolean (TRUE) indicating if a connection to ScienceBase can be established and if it is responding as expected. FALSE otherwise.

**Examples**

```r
#TRUE if all is well and SB can be contacted
sb_ping()
```

---

**session_details**  

*Get session info (deprecated)*

**Description**

Get the details associated with current ScienceBase user session.

**Usage**

```
session_details()
```

**Value**

list, if not logged in states that, but if logged in, user details
session_renew

Examples

## Not run:

session_details()

## End(Not run)

session_renew

Checks current session and re-authenticates if necessary

Description

Checks the state of your Sciencebase session, re-authenticates if the session is expired, and simply renews if the session is active.

Usage

session_renew(password, ..., username)

Arguments

password

The password to use, if needed, to renew the session.

...

Any additional parameters are currently ignored.

username

Optional. Used only to confirm that the current username is what you expect; if you want to switch usernames, use authenticate_sb() instead of this function.

Value

Returns the session object.

Examples

## Not run:

# an empty call is sufficient if the session is current,
# but will break if haven't been logged in before
session_renew()

# include a password if session may be expired
session_renew('newpass')

# optionally confirm the value of the current username
session_renew(username='olduser@usgs.gov', 'newpass')

## End(Not run)
**session_validate**  
*Validate sbtools session state*

**Description**

A session is considered valid if it is NULL or a true, non-expired SB session.

**Usage**

```r
session_validate()
```

**Details**

This function only operates on the active initialized session.

**Value**

TRUE/FALSE indicating if session is valid and can be used. Returns TRUE if session is NULL as well.

---

**set_endpoint**  
*Set SB endpoint*

**Description**

Sets the internal URLs used to either the production or development (beta) SB server. URLs are stored internally to the package.

**Usage**

```r
set_endpoint(endpoint = c("production", "development"))
```

**Arguments**

- `endpoint`  
  Indicate which SB endpoint you want to use: `c('production', 'development')`

**Author(s)**

Luke Winslow
Examples

```r
set_endpoint('prod')

# getting item from production SB servers
item_get('5060b03ae4b00fc20c4f3c8b')

set_endpoint('dev')
# getting item from beta SB servers
item_get('521e4686e4b051c878dc35d0')
```

---

**user_id** | *Get your parent ID*

Description

Required for creating items

Usage

```r
user_id(...)```

Arguments

... Additional parameters are passed on to `POST`

Value

A single character string, your user id

Examples

```r
## Not run:
user_id()

## End(Not run)```
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