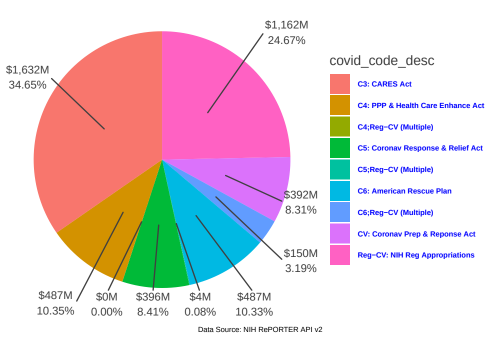
repoRter.nih: a convenient R interface to the NIH RePORTER Project API

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Legislative Source for NIH Covid Response Project Funding

Introduction

The US National Institute of Health (NIH) received funding of approximately \$42 billion in fiscal year 2022; \$31 billion (72%) of this was awarded by the NIH in the form of research grant funding to hospitals, medical colleges, non-profits, businesses, and other organizations based in the U.S. and abroad.¹ The NIH maintains a publicly available database called "RePORTER" to track this substantial flow of grant funding and makes it available to the public via a web-based query interface as well as an API.

"The NIH RePORTER APIs is designed to programmatically expose relevant scientific awards data from both NIH and non-NIH federal agencies for the consumption of project teams or external 3rd party applications to support reporting, data analysis, data integration or to satisfy other business needs as deemed pertinent." –NIH RePORTER v2 API Documentation

This data can have significant value for many audiences, including researchers, investors, industry, watchdogs/public advocates, and R users. But constructing queries and retrieving results programmatically involves some coding overhead which can be a challenge for those not familiar with RESTful APIs and JSON; it

 $^{^{1} \}rm https://nexus.od.nih.gov/all/2021/04/21/fy-2020-by-the-numbers-extramural-investments-in-research$

takes some effort even for those who are. The repoRter.nih package aims to simplify this task for the typical analyst scripting in R.

Getting Started

Installation

This package (latest stable release) can be installed from CRAN the usual way:

```
install.packages("repoRter.nih")
```

The current dev version can be installed from github, on the dev branch:

```
devtools::install_github('bikeactuary/repoRter.nih@dev')
```

I welcome R developers more capable than myself to collaborate on improving the source code, documentation, and unit testing in this package.

Basic Workflow

library(repoRter.nih)

The make_req() method is used to generate a valid JSON request object. The req can subsequently be passed to the RePORTER Project API and results retrieved via the get_nih_data() method.

Generating the request:

```
# all projects funded by the Paycheck Protection Act, Coronavirus Response and
# Relief Act, and American Rescue Plan, in fiscal year 2021
req <- make_req(criteria =</pre>
                  list(fiscal_years = 2021,
                        covid_response = c("C4", "C5", "C6")))
#> This is your JSON payload:
#>
   ſ
#>
       "criteria": {
#>
           "fiscal_years": [
#>
               2021
#>
           ],
#>
           "covid_response": [
#>
               "C4",
                "C5"
#>
                "C6"
#>
#>
           ],
#>
           "use_relevance": false,
           "include_active_projects": false,
#>
#>
           "exclude_subprojects": false,
#>
           "multi_pi_only": false,
           "newly_added_projects_only": false,
#>
#>
           "sub_project_only": false
#>
       },
#>
       "offset": 0,
       "limit": 500
#>
#> }
#>
#> If you receive a non-200 API response, compare this formatting (boxes, braces, quotes, etc.) to
    the 'Complete Payload' schema provided here:
#> https://api.reporter.nih.gov/?urls.primaryName=V2.0#/Search/post_v2_projects_search
```

Sending the request and retrieving results:

res <- get_nih_data(req)
#> Retrieving first page of results (up to 500 records)
class(res)
#> [1] "tbl_df" "tbl" "data.frame"

A tibble is returned containing 43 columns. This data is not flat - several columns are nested data.frames and lists (of variable length vectors and data.frames of varying height).

res %>% glimpse(width = getOption("cli.width")) #> Rows: 251 #> Columns: 44 <int> 10255113, 10425707, 10403857, 10258548, 10439178, 10446500, ~ #> \$ appl_id #> \$ subproject_id <int> 2021, 2021, 2021, 2021, 2021, 2021, 2021, 2021, 2021, 2021, 2021, #> \$ fiscal_year <chr> "3P20GM104417-07S1", "3P20GM104417-08S1", "3R01ES028615-07S1~ #> \$ project_num <chr> "GM104417", "GM104417", "ES028615", "ES028615", "DC019579", ~ #> \$ project_serial_num #> \$ organization <df[,17]> <data.frame[31 x 17]> <chr> "3", "3", "3", "3", "7", "3", "1", "3", "3", "1", "1", "~ #> \$ award_type <chr> "P20", "P20", "R01", "R01", "U01", "R01", "R01", "U01", "U19~ #> \$ activity_code #> \$ award_amount <int> 1115953, 681188, 300000, 1609765, 877287, 348242, 667277, 26~ <lp><lgl> TRUE, TRUE, TRUE, TRUE, TRUE, FALSE, TRUE, TRUE, FALSE~ #> \$ is_active #> \$ project_num_split <df[,7]> <data.frame[31 x 7]> #> \$ principal_investigators <list> [<data.frame[1 x 7]>], [<data.frame[1 x 7]>], [<data.frame[2-</pre> <chr> "ADAMS, ALEXANDRA K.", "ADAMS, ALEXANDRA K.", "AL-HENDY, ~ #> \$ contact_pi_name <list> [<data.frame[1 x 4]>], [<data.frame[1 x 4]>], [<data.frame[~ #> \$ program_officers <df[,3]> <data.frame[31 x 3]> #> \$ agency_ic_admin t> [<data.frame[1 x 5]>], [<data.frame[1 x 5]>], [<data.frame[1~ #> \$ agency_ic_fundings <chr> "MT-00", "MT-00", "IL-01", "IL-01", "MA-08", "MA-08", "MO-0~ #> \$ cong_dist #> \$ spending_categories <chr> "2020-11-17T12:11:00Z", "2021-09-01T12:09:00Z", "2020-11-11~ <chr> "2022-08-31T12:08:00Z", "2022-10-31T12:10:00Z", "2022-07-31T~ #> \$ project_start_date #> \$ project_end_date <df[,3]> <data.frame[31 x 3]> #> \$ organization_type <chr> "PA-20-135", "PAR-18-264", "PA-20-272", "PA-20-135", "RFA-OD~ #> \$ full_foa #> \$ full_study_section <df[,6]> <data.frame[31 x 6]> #> \$ award_notice_date <chr> "2020-11-17T12:11:00Z", "2021-09-21T12:09:00Z", "2021-08-31T~ **#> \$** is_new FALSE, FALS <chr> "RC", "RC", "RP", #> \$ mechanism_code_dc <chr> "P20GM104417", "P20GM104417", "R01ES028615", "R01ES028615", ~ #> \$ core_project_num <chr> "<Adult><21+ years old><Adult Human><adulthood><Affect><A~ #> \$ terms <chr> "2019-nCoV;Adult;Affect;Agricultural Workers;American Indian~ #> \$ pref_terms #> \$ abstract_text <chr> "Project Summary\nThe COVID-19 pandemic has disproportionate~ #> \$ project_title <chr> "Center for American Indian and Rural Health Equity", "Cente-<chr> "Project Narrative\nWorking with our Latino community partne" #> \$ phr_text <chr> "NIH", "NIH","","NIH","NIH",""NIH",""NIH",""NIH",""NIH",""NIH",""NIH",""N #> \$ agency_code <list> "C4", "C6", "C6", "C4", "C4", "C6", "C6", "C6", "C6", "C4", ~ #> \$ covid_response #> \$ arra_funded #> \$ budget_start <chr> "2020-11-17T12:11:00Z", "2021-09-01T12:09:00Z", "2021-09-01T~ #> \$ budget_end <chr> "2022-08-31T12:08:00Z", "2022-10-31T12:10:00Z", "2022-07-31T~ #> \$ cfda_code #> \$ funding_mechanism <chr> "Research Centers", "Research Centers", "Non-SBIR/STTR", "N~ #> \$ direct_cost_amt <int> 1006607, 616778, 297064, 1560464, 569403, 207287, 473684, 15~ #> \$ indirect_cost_amt <int> 109346, 64410, 2936, 49301, 307884, 140955, 193593, 116250, ~ #> \$ project_detail_url <chr> "https://reporter.nih.gov/project-details/10255113", "https:~ <chr> "2020-11-21T07:11:17Z", "2021-09-25T04:09:53Z", "2021-09-04T~ #> \$ date_added

Criteria-Field Translation

A dataset (nih_fields) is provided with this package to assist in translating between field names used in the payload criteria, column names in the return data, and field names used in the include_fields, exclude_fields, and sort_field arguments.

	("nih_fields") fields %>% print				
_	t A tibble: 43 x 5				
#>	payload_name	response_name	include_name	return_class	mod_ind
#>	<chr></chr>	<chr></chr>	<chr></chr>	< chr >	<int></int>
#>	1 appl_ids	appl_id	ApplId	integer	1
#>	2 <na></na>	subproject_id	SubprojectId	character	0
#>	3 fiscal_years	fiscal_year	FiscalYear	integer	1
#>	4 project_nums	project_num	ProjectNum	character	1
#>	5 serial_num	<pre>project_serial_num</pre>	ProjectSerialNum	character	1
#>	6 <na></na>	organization	Organization	data.frame	0
#>	7 award_types	award_type	AwardType	character	1
#>	8 activity_codes	activity_code	ActivityCode	character	1
#>	9 award_amount_range	award_amount	AwardAmount	integer	1
#> 1	0 include_active_projects	is_active	IsActive	logical	1
#> #	t with 33 more rows	_		0	

Some fields can not be used as filtering criteria - these will show NA in the payload_name column.

Generating Requests

Most of the detail (and function documentation) is around the many parameters available in RePORTER to filter/search project records. Let's get into some of the capabilities.

Default Request

If no arguments are supplied, the default behavior of make_req() is to generate a request for all projects funded in fiscal_years = lubridate::year(Sys.Date()). Limiting requests to a single year is often necessary (depending on additional filtering criteria used) due to a RePORTER restriction that a maximum of 10K records may be returned from any result set. There are currently ~2.6M projects in the database going back to fiscal year 1985, and each fiscal year tends to have 70-100K projects, so the 10K limit can be restrictive to the user wanting a broad search.

```
req <- make_req()</pre>
#> This is your JSON payload:
#> {
#>
        "criteria": {
#>
            "fiscal_years": [
#>
                2022
#>
            ],
            "use_relevance": false,
#>
#>
            "include_active_projects": false,
#>
            "exclude_subprojects": false,
#>
            "multi_pi_only": false,
            "newly_added_projects_only": false,
#>
#>
            "sub_project_only": false
#>
       },
       "offset": 0,
#>
#>
        "limit": 500
#> }
#>
```

```
#> If you receive a non-200 API response, compare this formatting (boxes, braces, quotes, etc.) to
    the 'Complete Payload' schema provided here:
#> https://api.reporter.nih.gov/?urls.primaryName=V2.0#/Search/post_v2_projects_search
```

The method prints a helpful message to the console in addition to returning the JSON. Set message = FALSE if you wish to suppress this message.

Limiting Data Retrieved

You can limit both the width and height of the result set retrieved from the API.

Fields

We probably will not need to fetch every field every time. The include_fields argument is provided to specify a limited set of fields to be returned. Alternatively, fields may be excluded using exclude_fields.

Records (projects)

This package provides the ability to retrieve only a limited number of result pages via the max_pages argument. This can be useful for developing/testing your queries (and for reducing time to render package documentation). Each page has a record count equal to limit - so setting max_pages = 5 with the default limit = 500 (the maximum permitted by RePORTER) in make_req() will result in up to 2,500 total records returned.

Ex. 1 - Limiting results and selecting fields

```
data("nih fields")
fields <- nih_fields %>%
      filter(response_name %in%
                                c("appl_id", "subproject_id", "project_title", "fiscal_year",
                                      "award_amount", "is_active", "project_start_date")) %>%
      pull(include_name)
req <- make_req(include_fields = fields,</pre>
                                              limit = 500,
                                             message = FALSE) # default
res <- get_nih_data(query = req,</pre>
                                                        max_pages = 1)
 #> Retrieving first page of results (up to 500 records)
 #> max_pages set to 1 by user. Result set contains 60 pages. Only partial results will be
             retrieved.
res %>% glimpse(width = getOption("cli.width"))
 #> Rows: 500
 #> Columns: 7
#> $ subproject_id
#> $ fiscal_year
#> $ award_amount
#> $ is contemport
 #> $ appl_id
                                                                   <int> 10464471, 10335890, 10376340, 10292904, 10388161, 10400390, 103941~
                                                                  nt <int> 67581, 244676, 387758, NA, 958638, 38717, NA, 384321, 47003, 70583~
<lgl> TRUE. TRUE. TRUE TRUE TRUE TRUE TRUE TRUE
                                                                  <lgl> TRUE, TR
 #> $ project_start_date <chr> "2022-05-01T12:05:00Z", "2020-02-28T12:02:00Z", "2019-04-01T12:04:~
 #> $ project_title  <chr>> "Elucidating novel molecular mechanisms of irisin-mediated effects~
```

Some Vanilla Criteria

Many criteria are passed as vectors within the **criteria** list argument. We will cover some of the most useful examples:

Ex. 2 - Organization search

We can refine our query results by providing filtering criteria to make_req(), and by extension to the API. Suppose we want all currently active projects, funded in fiscal years 2017 through 2021, with a specific organization in mind (though we don't know exactly how its name will appear in RePORTER):

```
req <- make_req(criteria =
    list(
        fiscal_years = 2010:2011,
        include_active_projects = TRUE,
        org_names = c("Yale", "New Haven")
        ),
        include_fields = c("Organization", "FiscalYear", "AwardAmount"),
        message = FALSE)</pre>
```

Here we are asking for any orgs containing the strings "yale" or "new haven" (ignoring case) - there are implied wildcards on either end of the strings we provide. This is the same as org_name LIKE '%yale%' OR org_name LIKE '%new haven%' in a SQL WHERE clause.

```
res <- get_nih_data(req, max_pages = 1)
#> Retrieving first page of results (up to 500 records)
#> max_pages set to 1 by user. Result set contains 8 pages. Only partial results will be retrieved.
res %>% glimpse(width = getOption("cli.width"))
#> Rows: 500
#> Columns: 3
#> $ fiscal_year <int> 2022, 2022, 2022, 2022, 2022, 2022, 2022, 2022, 2022, 2022, 2022, 2022, 2022, 2022, ~
#> $ organization <df[,17]> <data.frame[31 x 17]>
#> $ award_amount <int> 592057, 165660, 4777718, 460626, 30891, 770500, 413843, 46753, 548969~
```

Notice the column organization is a nested data frame - it has 17 columns and always a single record. Setting flatten_result = TRUE in the call to get_nih_data() will flatten all such return fields, prefixing the original field name and returning with clean names (see janitor::clean_names()).

```
res <- get_nih_data(req,</pre>
                                                            \max_{pages} = 1,
                                                            flatten_result = TRUE)
#> Retrieving first page of results (up to 500 records)
#> max_pages set to 1 by user. Result set contains 8 pages. Only partial results will be retrieved.
res %>% glimpse(width = getOption("cli.width"))
#> Rows: 500
#> Columns: 19
#> $ fiscal_year
                                                                                                            <int> 2022, 2022, 2022, 2022, 2022, 2022, 2022, 2022, 2022, ~
                                                                                                            <int> 592057, 165660, 477718, 460626, 30891, 770500, 413843,~
#> $ award amount
                                                                                                            <chr> "YALE UNIVERSITY", "YALE UNIVERSITY", "YALE UNIVERSITY~
#> $ organization_org_name
#> $ organization_city
                                                                                                            #> $ organization_country
                                                                                                            <chr> "NEW HAVEN", "NEW HAVEN", "NEW HAVEN", "NEW HAVEN", "N~
#> $ organization_org_city
                                                                                                            <chr> "UNITED STATES", "UNITED STATES", "UNITED STATES", "UN-
#> $ organization_org_country
                                                                                                            <chr> "CT", 
#> $ organization_org_state
                                                                                                            #> $ organization_org_state_name
#> $ organization_dept_type
                                                                                                            <chr> "RADIATION-DIAGNOSTIC/ONCOLOGY", "INTERNAL MEDICINE/ME~
<chr> "043207562", "043207562", "043207562", "043207562", "0-
#> $ organization_org_duns
                                                                                                            <chr> "FL6GV84CKN57", "FL6GV84CKN57", "FL6GV84CKN57", "FL6GV~
#> $ organization_org_ueis
                                                                                                            <chr> "043207562", "043207562", "043207562", "043207562", "0~
#> $ organization_primary_duns
                                                                                                            <chr> "FL6GV84CKN57", "FL6GV84CKN57", "FL6GV84CKN57", "FL6GV~
#> $ organization_primary_uei
                                                                                                            <chr> "US", "US", "US", "US", "US", "US", "US", "US", "US", ~
#> $ organization_org_fips
#> $ organization_org_ipf_code <chr> "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "9420201", "94200", "94200", "94200", "94200", "94200", "9420", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "94200", "9400", "9400", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "94000", "940000", "94000", "940000",","%
```

```
#> $ organization_org_zipcode <chr> "065208327", "065208327", "065208327", "065208327", "065208327", "0~
#> $ organization_external_org_id <int> 9420201, 9420201, 9420201, 9420201, 9420201, 9420201, ~
```

Most users will prefer the flattened format above. It looks like Yale is busy, but it is not the only org matching our search.

The org_names_exact_match criteria can be used as an alternative when we know the exact org name as it appears in RePORTER, if we want only that org's projects returned.

Ex. 3 - Geographic search

We can also filter projects by the geographic location (country/state/city) of the applicant organization.

```
## A valid request but probably not what we want
req <- make_req(criteria =</pre>
                  list(
                     fiscal_years = 2010:2011,
                    include_active_projects = TRUE,
                    org_cities = "New Haven",
                    org_states = "WY"
                  ),
                include_fields = c("Organization", "FiscalYear", "AwardAmount"),
                message = FALSE ## suppress printed message
)
res <- get_nih_data(req,</pre>
                    max_pages = 5,
                    flatten_result = TRUE)
#> Retrieving first page of results (up to 500 records)
#> Done - 0 records returned. Try a different search criteria.
```

Multiple criteria are usually connected by logical "AND" - there are no orgs based in the city of New Haven in Wyoming state (because it doesn't exist.)

<pre>res %>% glimpse(width = getOption("cli.width"))</pre>				
#> Rows: 97	> Rows: 97			
#> Columns: 20				
<pre>#> \$ appl_id</pre>	<int> 8884461, 8898483, 10398032, 10201479, 10216275, 103980~</int>			
#> \$ fiscal_year	<int> 2015, 2015, 2022, 2021, 2021, 2022, 2015, 2015, 2021, ~</int>			
<pre>#> \$ award_amount</pre>	<int> 4957554, 3521553, 3418046, 2638712, 2004078, 1668383, ~</int>			
<pre>#> \$ organization_org_name</pre>	<chr> "WYOMING STATE DEPARTMENT OF HEALTH", "UNIVERSITY OF W~</chr>			
<pre>#> \$ organization_city</pre>	<lgl> NA, NA, NA, NA, NA, NA, NA, NA, NA, NA,</lgl>			
<pre>#> \$ organization_country</pre>	<lgl> NA, NA, NA, NA, NA, NA, NA, NA, NA, NA,</lgl>			
<pre>#> \$ organization_org_city</pre>	<chr> "CHEYENNE", "LARAMIE", "LARAMIE", "LARAMIE", "LARAMIE"~</chr>			
<pre>#> \$ organization_org_country</pre>	<chr> "UNITED STATES", "UNITED STATES", "UNITED STATES", "UN~</chr>			
<pre>#> \$ organization_org_state</pre>	<chr> "WY", "WY", "WY", "WY", "WY", "WY", "WY", "WY", "WY", ~</chr>			
<pre>#> \$ organization_org_state_name</pre>	<lgl> NA, NA, NA, NA, NA, NA, NA, NA, NA, NA,</lgl>			
<pre>#> \$ organization_dept_type</pre>	<chr> NA, "PHARMACOLOGY", "PHARMACOLOGY", "VETERINARY SCIENC~</chr>			
<pre>#> \$ organization_fips_country_code</pre>	<lgl> NA, NA, NA, NA, NA, NA, NA, NA, NA, NA,</lgl>			
<pre>#> \$ organization_org_duns</pre>	<chr> "809915796", "069690956", "069690956", "069690956", "0~</chr>			
<pre>#> \$ organization_org_ueis</pre>	<chr> "JP1QRJYYJG73", "FDR5YF2K32X5", "FDR5YF2K32X5", "FDR5Y~</chr>			
<pre>#> \$ organization_primary_duns</pre>	<chr> "809915796", "069690956", "069690956", "069690956", "0~</chr>			
<pre>#> \$ organization_primary_uei</pre>	<chr> "JP1QRJYYJG73", "FDR5YF2K32X5", "FDR5YF2K32X5", "FDR5Y~</chr>			
<pre>#> \$ organization_org_fips</pre>	<chr> "US", "US", "US", "US", "US", "US", "US", "US", "US", ~</chr>			
<pre>#> \$ organization_org_ipf_code</pre>	<chr> "9408801", "9412601", "9412601", "9412601", "9412601",~</chr>			
<pre>#> \$ organization_org_zipcode</pre>	<chr> "820020001", "820712000", "820712000", "820712000", "8~</chr>			
<pre>#> \$ organization_external_org_id</pre>	<int> 9408801, 9412601, 9412601, 9412601, 9412601, 9412601, ~</int>			

Why are there projects from more recent years than 2015? Because the include_active_projects flag adds in active projects that match all criteria aside from fiscal_years (this appears to be the intended behavior by RePORTER).

Ex. 3 - Coronavirus/Covid-19 research

We already provided one example of this search criteria above. Let's mix it up and request all Covid response projects.

```
## all projects funded by the Paycheck Protection Act, Coronavirus Response and Relief Act,
## and American Rescue Plan, over all years
req <- make_req(criteria =</pre>
                  list(covid_response = c("All")),
                include_fields = nih_fields %>%
                  filter(payload_name %in% c("award_amount_range", "covid_response"))
                %>% pull(include_name))
#> This is your JSON payload:
#> {
       "criteria": {
#>
           "covid_response": [
#>
               "All"
#>
#>
           ],
           "use_relevance": false,
#>
#>
           "include_active_projects": false,
           "exclude_subprojects": false,
#>
#>
           "multi_pi_only": false,
#>
           "newly_added_projects_only": false,
#>
           "sub_project_only": false
#>
       },
       "include_fields": [
#>
#>
           "AwardAmount",
#>
           "CovidResponse"
#>
       ],
       "offset": 0,
#>
       "limit": 500
#>
```

```
#> }
#>
#>
#>
#>
If you receive a non-200 API response, compare this formatting (boxes, braces, quotes, etc.) to
    the 'Complete Payload' schema provided here:
#> https://api.reporter.nih.gov/?urls.primaryName=V2.0#/Search/post_v2_projects_search
res <- get_nih_data(req)
#> Retrieving first page of results (up to 500 records)
#> Retrieving results 501 to 1000 of 2698
#> Retrieving results 1001 to 1500 of 2698
#> Retrieving results 1501 to 2000 of 2698
#> Retrieving results 2001 to 2500 of 2698
#> Retrieving results 2001 to 2500 of 2698
#> Retrieving results 2001 to 2500 of 2698
#> Retrieving results 2501 to 2698 of 2698
```

Let's inspect the result:

```
res$covid_response %>% class()
#> [1] "list"
res$covid_response[[1]]
#> [1] "Req-CV"
```

covid_response is a nested list (with character vectors of variable length) within the return tibble. We can
use flatten_result = TRUE here - elements of each vector will be collapsed to a single string delimited by ";",
massaging the list to a single character vector.

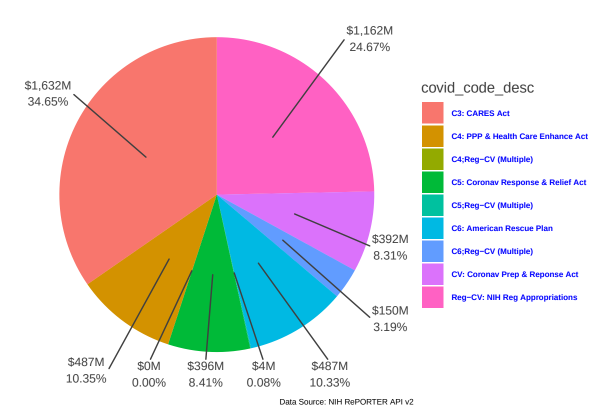
unique(res\$covid_response)

#>	[1]	"Reg-CV"	"CV"	"C3"	"C4"	"C6"	"C6;Reg-CV" "C5"
#>	[8]	"C5;Reg-CV"	"C4;Reg-CV"				

Some projects are being funded from multiple sources. Summarizing all Covid-related project awards:

```
library(ggplot2)
```

```
accuracy = 1,
                                    suffix = "M").
                             "\n", percent(prop, accuracy = .01)),
                    y = ypos),
                show.legend = FALSE,
                nudge_x = .8,
                size = 3, color = "grey25") +
coord_polar(theta ="y") +
theme_void() +
theme(legend.position = "right",
      legend.title = element_text(colour = "grey25"),
      legend.text = element_text(colour="blue", size=6,
                                 face="bold"),
      plot.title = element_text(color = "grey25"),
      plot.caption = element_text(size = 6)) +
labs(caption = "Data Source: NIH RePORTER API v2") +
ggtitle("Legislative Source for NIH Covid Response Project Funding")
```



Legislative Source for NIH Covid Response Project Funding

A second dataset is provided to translate the covid_response codes; it includes both the long-form and a shorter version of the source name.

data("co	data("covid_response_codes")						
covid_re	covid_response_codes %>% print						
#> # A t	#> # A tibble: 6 x 3						
#> cov	id_response funding_source	$fund_src$					
#> <ch< td=""><td>r> <chr></chr></td><td><chr></chr></td></ch<>	r> <chr></chr>	<chr></chr>					
#> 1 Reg	-CV NIH regular appropriations funding	NIH Reg~					
#> 2 CV	Coronavirus Preparedness and Response Supplem	nental Appropriations Ac~ Coronav~					

#	> 3 C3	CARES Act (Coronavirus Aid, Relief, and Economic Security Act), 2020	CARES A~
#	\$> 4 C4	Paycheck Protection Program and Health Care Enhancement Act, 2020	PPP & H~
#	\$> 5 C5	Coronavirus Response and Relief Supplemental Appropriations Act, 2021	Coronav~
#	!> 6 C6	American Rescue Plan Act of 2021	America~

Some Rocky Road Criteria

Other criteria provide search and filtering capability on many of the nested data elements. These criteria are passed as lists and must include a value for each of the named elements within.

Ex. 4 - Principal Investigator/Officer name search

The pi_names and po_names criteria allow the user to search for projects based on the first and last names of Principal Investigators and Principal Officers assigned. Each of these criteria must be provided as a list with three named character vector elements: first_name, last_name, and any_name. Even if you only want to search on one of these name fields, you must provide the remaining elements as an empty string. We will demonstrate with a search on PI name:

```
## projects funded in 2021 where the principal investigator first name
## is "Michael" or begins with "Jo"
req <- make_req(criteria =</pre>
                  list(fiscal_years = 2021,
                       pi_names = list(first_name = c("Michael", "Jo*"),
                                        last_name = c(""), # must specify all pi_names elements
    always
                                        any_name = character(1))),
                include_fields = nih_fields %>%
                  filter(payload_name == "pi_names") %>%
                  pull(include_name),
                message = FALSE)
res <- get_nih_data(req,</pre>
                    max_pages = 1,
                    flatten_result = TRUE)
#> Retrieving first page of results (up to 500 records)
#> max_pages set to 1 by user. Result set contains 13 pages. Only partial results will be
    retrieved.
res %>% glimpse(width = getOption("cli.width"))
#> Rows: 500
#> Columns: 2
#> $ principal_investigators <list> [<data.frame[3 x 7]>], [<data.frame[2 x 7]>], [<data.frame[2</pre>
                              <chr> "WU, JOSEPH C.", "DIPERSIO, C. MICHAEL ", "ROUT, MICHAEL P", ~
#> $ contact_pi_name
```

Here we searched for any projects with a PI first-named "Michael" or beginning with "Jo" - the "*" is a wildcard operator.

Note that the first column in the return is a list of data frames of *variable height* (not a nested data.frame) - we leave such returned elements to the user to handle extraction/formatting - flattening is only performed for lists of atomic vectors and nested data frames.

Ex. 5 - Advanced text search

RePORTER allows users to search the project title, abstract, and tags for specific terms or phrases. You can access this capability with the advanced_text_search criteria - a named list with three elements:

- operator may be either "and", "or", or "advanced" and/or will specify the logical operator connecting multiple search terms. "advanced" allows the user to pass a boolean search string directly;
- search_field can be any or multiple of "terms", "abstract", "projecttitle." To search all items, specify "all" or "" (a length 1 vector with an empty string);
- search_text may be either a length 1 character vector of space-delimited search terms (when using "and" or "or" for the operator argument the logical operator is inserted between all search terms); or it may be a boolean search string (when specifying "advanced" for the operator argument).

```
## using advanced_text_search with boolean search string
req <- make_req(criteria =</pre>
                  list(advanced_text_search =
                         list(operator = "advanced",
                               search_field = c("terms", "abstract"),
                               search_text = "(head AND trauma) OR \"brain damage\" AND NOT
    \"psychological\"")),
                include_fields = c("ProjectTitle", "AbstractText", "Terms") )
#> This is your JSON payload:
#> {
#>
       "criteria": {
#>
           "advanced_text_search": {
               "operator": "advanced",
#>
               "search_field": "terms, abstract",
#>
               "search_text": "(head AND trauma) OR \"brain damage\" AND NOT \"psychological\""
#>
#>
           },
#>
           "use_relevance": false,
#>
           "include_active_projects": false,
#>
           "exclude_subprojects": false,
           "multi_pi_only": false,
#>
#>
           "newly_added_projects_only": false,
#>
           "sub_project_only": false
#>
       }.
       "include_fields": [
#>
#>
           "ProjectTitle",
           "AbstractText",
#>
           "Terms"
#>
#>
       ],
#>
       "offset": 0,
       "limit": 500
#>
#> }
#>
#> If you receive a non-200 API response, compare this formatting (boxes, braces, quotes, etc.) to
    the 'Complete Payload' schema provided here:
#> https://api.reporter.nih.gov/?urls.primaryName=V2.0#/Search/post_v2_projects_search
res <- get_nih_data(req, max_pages = 1)</pre>
#> Retrieving first page of results (up to 500 records)
#> max_pages set to 1 by user. Result set contains 38 pages. Only partial results will be
    retrieved.
```

Let's inspect the fields we searched from one of these results:

```
## substr to avoid LaTeX error exceeding char limit
one_rec %>% pull(abstract_text) %>% substr(1, 85) %>% print
#> [1] "ABSTRACT/PROJECT SUMMARY Linked VA Merit Overall Research Strategy: Traumatic brain i"
```

```
one_rec %>% pull(terms) %>% substr(1, 85) %>% print
#> [1] "<Affect><After Care><After-Treatment><post treatment><Aftercare><ages><Age><AD dement"</pre>
```

Large Result Sets

The RePORTER API provides no direct way to obtain complete result sets when searches yield over 10,000 records. get_nih_data() provides the return_meta argument which is defaulted to FALSE. When set to TRUE and combined with a little programming, you can easily obtain full result sets well beyond the 10K limit. One approach may be the following:

- 1. Obtain a sample from your full result set by making the query you desire and calling get_nih_data()
 with max_pages = 1 (or some small number of pages); also set return_meta = TRUE in order to determine
 the total number of records in the full result set
- 2. Calculate quantiles for the sample distribution of a column of your choice (e.g. award_amount)
- Set the # of quantiles such that you can confidently infer that the number of records within each quantile range will contain <10K records within the full result set
- 3. Iterate over your quantiles making separate requests, passing the endpoints of each quantile to award_amount_range criteria
- Wait until the end to flatten the combined results since some columns may flatten differently on smaller individual result sets, causing problems in combining them after flattening
- 4. Bind your list of results together
- 5. Flatten the complete result set, if desired

Below is an implementation of the above logic:

```
all res <- list()
for(y in 2017:2021) { ## five years to loop over, each year is ~80K records
  ## We only need the AwardAmount for quantiles
 req_sample <- make_req(criteria = list(fiscal_years = y),</pre>
                         include_fields = "AwardAmount")
  ## get a sample of the result set - 1000 records should be enough
  ## return the metadata
 res_sample <- get_nih_data(req_sample, max_pages = 2, return_meta = TRUE)</pre>
 paste0("There are ", res_sample$meta$total, " results for fiscal year ", y) %>%
    print()
  ## deciles of award amount - each decile should contain ~7,314.2 records, approximately
  qtiles <- res_sample$records %>% pull(award_amount) %>% quantile(na.rm = TRUE, probs = seq(.1,
    1, .1))
  ## list for qtile results (full year)
  this_res <- list()</pre>
  ## for each gtile
  for (i in 1:length(qtiles)) {
    if (i == 1) {
      award_min <- 0
   } else {
      award_min <- ceiling(qtiles[i-1])+.01</pre>
```

```
}
    if (i == length(qtiles)) {
      award_max <- 1e9 ## arbitrarily huge</pre>
    } else {
      award_max <- ceiling(qtiles[i])</pre>
    }
    req <- make_req(criteria = list(fiscal_years = y,</pre>
                                      award amount range = list(min amount = award min,
                                                                 max_amount = award_max)))
    ## result set for quantile
    this_res[[i]] <- get_nih_data(req, flatten_result = FALSE)</pre>
  }
  ## list of result sets for each year
 yr_res[[y %>% as.character()]] <- this_res</pre>
}
## shape it up
all_res <- unlist(yr_res, recursive = FALSE) %>%
  bind_rows() %>%
  flatten(recursive = FALSE) %>%
  clean_names()
## pull out everything that is flat
flat_columns <- all_res %>%
  select_if(is.atomic)
## everything that isn't
annoying_columns <- all_res %>%
  select_if(!is.atomic)
```

Note that using award_amount for this purpose will omit records with missing values. If you need these included, you may consider similar logic applied to an alternative field such as award_notice_date.

Additional Resources

- The **RePORTER** web interface and official API documentation are useful for getting familiar with available search parameters
- ... and the homepage with further examples/documentation is here
- Information on NIH study sections, IRGs, etc. is here
- h/t to Chris whose code on github was all I could find existing in R and served as a starting point for this work

Session Information

The version number of R and packages loaded for generating the vignette were:

```
sessionInfo()
#> R version 4.1.0 (2021-05-18)
#> Platform: x86_64-pc-linux-gnu (64-bit)
#> Running under: Ubuntu 20.04.2 LTS
#>
#> Matrix products: default
#> BLAS: /usr/lib/x86_64-linux-gnu/blas/libblas.so.3.9.0
#> LAPACK: /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.9.0
#>
#> locale:
```

```
#> [1] LC_CTYPE=C.UTF-8 LC_NUMERIC=C LC_TIME=C.UTF-8
#> [4] LC_COLLATE=C
                          LC_MONETARY=C.UTF-8
                                               LC_MESSAGES=C.UTF-8
#> [7] LC_PAPER=C.UTF-8
                         LC_NAME=C
                                               LC_ADDRESS=C
#> [10] LC_TELEPHONE=C
                          LC_MEASUREMENT=C.UTF-8 LC_IDENTIFICATION=C
#>
#> attached base packages:
#> [1] stats graphics grDevices utils
                                        datasets methods base
#>
#> other attached packages:
#> [1] repoRter.nih_0.1.3 tufte_0.12
                                       scales_1.2.0
                                                        dplyr_1.0.7
#> [5] ggrepel_0.9.1 ggplot2_3.3.6
                                       tibble_{3.1.7}
#>
#> loaded via a namespace (and not attached):
#> [1] Rcpp_1.0.7 highr_0.9 pillar_1.7.0
                                                     compiler_4.1.0
                                                                    tools_4.1.0
                    jsonlite_1.7.2
#> [6] digest_0.6.29
                                      lubridate_1.7.10 evaluate_0.15
                                                                      lifecycle_1.0.1
#> [11] gtable_0.3.0
                     pkgconfig_2.0.3 rlang_1.0.2 cli_3.3.0
                                                                      DBI_1.1.1
#> [16] rstudioapi_0.13 curl_4.3.2
                                      yaml_2.2.1
                                                     xfun_0.24
                                                                      withr_2.5.0
#> [21] stringr_1.4.0 httr_1.4.2
                                      knitr_1.33
                                                      janitor_2.1.0 generics_0.1.0
#> [26] vctrs_0.4.1
                     grid_4.1.0
                                      tidyselect_1.1.1 glue_1.6.2
                                                                      snakecase_0.11.0
                     fansi_1.0.3
#> [31] R6_2.5.1
                                      rmarkdown_2.11 farver_2.1.0
                                                                      purrr_0.3.4
#> [36] magrittr_2.0.3 ellipsis_0.3.2
                                      htmltools_0.5.1.1 assertthat_0.2.1 colorspace_2.0-2
#> [41] labeling_0.4.2 utf8_1.2.2
                                      stringi_1.7.3 munsell_0.5.0 crayon_1.4.1
```