

Package ‘biocompute’

May 3, 2022

Type Package

Title Create and Manipulate BioCompute Objects

Version 1.1.1

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Description Tools to create, validate, and export BioCompute Objects described in King et al. (2019) <[doi:10.17605/osf.io/h59uh](https://doi.org/10.17605/osf.io/h59uh)>. Users can encode information in data frames, and compose BioCompute Objects from the domains defined by the standard. A checksum validator and a JSON schema validator are provided. This package also supports exporting BioCompute Objects as JSON, PDF, HTML, or 'Word' documents, and exporting to cloud-based platforms.

License AGPL-3

VignetteBuilder knitr

URL <https://sbg.github.io/biocompute/>,
<https://github.com/sbg/biocompute>

BugReports <https://github.com/sbg/biocompute/issues>

Encoding UTF-8

Imports methods, jsonlite, yaml, digest, uuid, jsonvalidate, httr,
curl, crayon, cli, stringr, magrittr, rmarkdown

Suggests knitr

RoxygenNote 7.1.2

NeedsCompilation no

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Repository CRAN

Date/Publication 2022-05-03 07:00:27 UTC

R topics documented:

compose_description_v1.4.2	2
compose_error_v1.4.2	5
compose_execution_v1.4.2	6
compose_extension_v1.4.2	8
compose_fhir_v1.4.2	9
compose_io_v1.4.2	10
compose_parametric_v1.4.2	11
compose_provenance_v1.4.2	12
compose_scm_v1.4.2	14
compose_tlf_v1.4.2	15
compose_usability_v1.4.2	16
compose_v1.4.2	17
convert_json	18
convert_yaml	19
export_html	20
export_json	20
export_pdf	21
export_sevenbridges	22
export_word	23
generate_example	24
generate_id	24
is_bco	25
is_domain	25
read_bco	26
validate_checksum_v1.4.2	26
validate_schema_v1.4.2	27
versions	28
Index	29

compose_description_v1.4.2

Compose BioCompute Object - Description Domain (v1.4.2)

Description

Compose BioCompute Object - Description Domain (v1.4.2)

Usage

```
compose_description_v1.4.2(
  keywords = NULL,
  xref = NULL,
  platform = list("Seven Bridges Platform"),
  pipeline_meta = NULL,
  pipeline_prerequisite = NULL,
```

```

    pipeline_input = NULL,
    pipeline_output = NULL
  )

compose_description(
  keywords = NULL,
  xref = NULL,
  platform = list("Seven Bridges Platform"),
  pipeline_meta = NULL,
  pipeline_prerequisite = NULL,
  pipeline_input = NULL,
  pipeline_output = NULL
)

```

Arguments

keywords	Character vector. A list of keywords to aid in searchability and description of the experiment.
xref	Data frame. A list of the databases and/or ontology IDs that are cross-referenced in the BCO.
platform	Character string or list. Reference to a particular deployment of an existing platform where this BCO can be reproduced.
pipeline_meta	Data frame. Pipeline metadata. Variables include step_number, name, description, and version.
pipeline_prerequisite	Data frame. Packages or prerequisites for running the tools used. Variables include step_number, name, uri, and access_time.
pipeline_input	Data frame. Input files for the tools. Variables include step_number, uri, and access_time.
pipeline_output	Data frame. Output files for the tools. Variables include step_number, uri, and access_time.

Value

A list of class `bco.domain`

Examples

```

keywords <- c("HCV1a", "Ledipasvir", "antiviral resistance", "SNP", "amino acid substitutions")
xref <- data.frame(
  "namespace" = c("pubchem.compound", "pubmed", "so", "taxonomy"),
  "name" = c("PubChem-compound", "PubMed", "Sequence Ontology", "Taxonomy"),
  "ids" = I(list(
    "67505836",
    "26508693",
    c("SO:000002", "SO:0000694", "SO:0000667", "SO:0000045"),
    "31646"
  )),
)

```

```

"access_time" = c(
  as.POSIXct("2017-01-20T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST"),
  as.POSIXct("2017-01-21T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST"),
  as.POSIXct("2017-01-22T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST"),
  as.POSIXct("2017-01-23T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST")
),
stringsAsFactors = FALSE
)

platform <- "Seven Bridges Platform"

pipeline_meta <- data.frame(
  "step_number" = c("1"),
  "name" = c("HIVE-hexagon"),
  "description" = c("Alignment of reads to a set of references"),
  "version" = c("1.3"),
  stringsAsFactors = FALSE
)

pipeline_prerequisite <- data.frame(
  "step_number" = rep("1", 5),
  "name" = c(
    "Hepatitis C virus genotype 1",
    "Hepatitis C virus type 1b complete genome",
    "Hepatitis C virus (isolate JFH-1) genomic RNA",
    "Hepatitis C virus clone J8CF, complete genome",
    "Hepatitis C virus S52 polyprotein gene"
  ),
  "uri" = c(
    "https://www.ncbi.nlm.nih.gov/nuccore/22129792",
    "https://www.ncbi.nlm.nih.gov/nuccore/5420376",
    "https://www.ncbi.nlm.nih.gov/nuccore/13122261",
    "https://www.ncbi.nlm.nih.gov/nuccore/386646758",
    "https://www.ncbi.nlm.nih.gov/nuccore/295311559"
  ),
  "access_time" = c(
    as.POSIXct("2017-01-24T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST"),
    as.POSIXct("2017-01-24T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST"),
    as.POSIXct("2017-01-24T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST"),
    as.POSIXct("2017-01-24T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST"),
    as.POSIXct("2017-01-24T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST")
  ),
  stringsAsFactors = FALSE
)

pipeline_input <- data.frame(
  "step_number" = rep("1", 2),
  "uri" = c(
    "https://example.com/dna.cgi?cmd=objFile&ids=514683",
    "https://example.com/dna.cgi?cmd=objFile&ids=514682"
  ),
  "access_time" = c(
    as.POSIXct("2017-01-24T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST"),

```

```

    as.POSIXct("2017-01-24T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST")
  ),
  stringsAsFactors = FALSE
)

pipeline_output <- data.frame(
  "step_number" = rep("1", 2),
  "uri" = c(
    "https://example.com/data/514769/allCount-aligned.csv",
    "https://example.com/data/514801/SNPPProfile*.csv"
  ),
  "access_time" = c(
    as.POSIXct("2017-01-24T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST"),
    as.POSIXct("2017-01-24T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST")
  ),
  stringsAsFactors = FALSE
)

compose_description(
  keywords, xref, platform,
  pipeline_meta, pipeline_prerequisite, pipeline_input, pipeline_output
) %>% convert_json()

```

compose_error_v1.4.2 *Compose BioCompute Object - Error Domain (v1.4.2)*

Description

The error domain can be used to determine what range of input returns outputs that are within the tolerance level defined in this subdomain and therefore can be used to optimize algorithm (**domain definition**).

Usage

```
compose_error_v1.4.2(empirical = NULL, algorithmic = NULL)
```

```
compose_error(empirical = NULL, algorithmic = NULL)
```

Arguments

empirical	Data frame. Variables include key and value. Each row is one item in the empirical error subdomain.
algorithmic	Data frame. Variables include key and value. Each row is one item in the algorithmic subdomain.

Value

A list of class `bco.domain`

Examples

```

empirical <- data.frame(
  "key" = c("false_negative_alignment_hits", "false_discovery"),
  "value" = c("<0.0010", "<0.05"),
  stringsAsFactors = FALSE
)

algorithmic <- data.frame(
  "key" = c("false_positive_mutation_calls", "false_discovery"),
  "value" = c("<0.00005", "0.005"),
  stringsAsFactors = FALSE
)

compose_error(empirical, algorithmic) %>% convert_json()

```

compose_execution_v1.4.2

Compose BioCompute Object - Execution Domain (v1.4.2)

Description

Compose BioCompute Object - Execution Domain (v1.4.2)

Usage

```

compose_execution_v1.4.2(
  script = NULL,
  script_driver = NULL,
  software_prerequisites = NULL,
  external_data_endpoints = NULL,
  environment_variables = NULL
)

compose_execution(
  script = NULL,
  script_driver = NULL,
  software_prerequisites = NULL,
  external_data_endpoints = NULL,
  environment_variables = NULL
)

```

Arguments

script	Character string or list. Points to internal or external references to an object that was used to perform computations for this BCO instance.
script_driver	Character string. Indicate what kind of executable can be launched in order to perform a sequence of commands described in the script in order to run the pipeline.

software_prerequisites

Data frame. The minimal necessary prerequisites, library, and tool versions needed to successfully run the script to produce BCO. Variables include name, version, uri, access_time, and sha1_chksum. Each row is one item in the output subdomain.

external_data_endpoints

Data frame. The minimal necessary domain-specific external data source access to successfully run the script to produce the BCO. Variables include mediatype, name, and url. Each row is one item in the output subdomain.

environment_variables

Data frame. Key-value pairs useful to configure the execution environment on the target platform. Variables include key and value.

Value

A list of class `bco.domain`

Examples

```
script <- "https://example.com/workflows/antiviral_resistance_detection_hive.py"
script_driver <- "shell"
software_prerequisites <- data.frame(
  "name" = c("HIVE-hexagon", "HIVE-heptagon"),
  "version" = c("babajanian.1", "albinoni.2"),
  "uri" = c(
    "https://example.com/dna.cgi?cmd=dna-hexagon&cmdMode=-",
    "https://example.com/dna.cgi?cmd=dna-heptagon&cmdMode=-"
  ),
  "access_time" = c(
    as.POSIXct("2017-01-24T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST"),
    as.POSIXct("2017-01-24T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST")
  ),
  "sha1_chksum" = c("d60f506cddac09e9e816531e7905ca1ca6641e3c", NA),
  stringsAsFactors = FALSE
)
external_data_endpoints <- data.frame(
  "name" = c("generic name", "access to ftp server", "access to e-utils web service"),
  "url" = c(
    "protocol://domain:port/application/path",
    "ftp://data.example.com:21/",
    "https://eutils.ncbi.nlm.nih.gov/entrez/eutils"
  ),
  stringsAsFactors = FALSE
)
environment_variables <- data.frame(
  "key" = c("HOSTTYPE", "EDITOR"),
  "value" = c("x86_64-linux", "vim")
)

compose_execution(
  script, script_driver, software_prerequisites, external_data_endpoints, environment_variables
) %>% convert_json()
```

`compose_extension_v1.4.2`*Compose BioCompute Object - Extension Domain (v1.4.2)*

Description

Compose BioCompute Object - Extension Domain (v1.4.2)

Usage

```
compose_extension_v1.4.2(fhir = NULL, scm = NULL)
```

```
compose_extension(fhir = NULL, scm = NULL)
```

Arguments

<code>fhir</code>	FHIR extension domain composed by compose_fhir .
<code>scm</code>	SCM extension domain composed by compose_scm .

Value

A list of class `bco.domain`

Examples

```
fhir_endpoint <- "https://fhirtest.uhn.ca/baseDstu3"
fhir_version <- "3"
fhir_resources <- data.frame(
  "id" = c("21376", "6288583", "25544", "92440", "4588936"),
  "resource" = c(
    "Sequence", "DiagnosticReport", "ProcedureRequest",
    "Observation", "FamilyMemberHistory"
  ),
  stringsAsFactors = FALSE
)

fhir <- compose_fhir(fhir_endpoint, fhir_version, fhir_resources)

scm_repository <- "https://github.com/example/repo"
scm_type <- "git"
scm_commit <- "c9ffea0b60fa3bcf8e138af7c99ca141a6b8fb21"
scm_path <- "workflow/hive-viral-mutation-detection.cwl"
scm_preview <- "https://github.com/example/repo/blob/master/mutation-detection.cwl"

scm <- compose_scm(scm_repository, scm_type, scm_commit, scm_path, scm_preview)

compose_extension(fhir, scm) %>% convert_json()
```

compose_fhir_v1.4.2 *Compose BioCompute Object - FHIR Extension (v1.4.2)*

Description

Compose BioCompute Object - FHIR Extension (v1.4.2)

Usage

```
compose_fhir_v1.4.2(endpoint = NULL, version = NULL, resources = NULL)
```

```
compose_fhir(endpoint = NULL, version = NULL, resources = NULL)
```

Arguments

endpoint	Character string. The URL of the endpoint of the FHIR server containing the resource.
version	Character string. The FHIR version used.
resources	Data frame with two variables: id and resource. Each row is one item of resources to fetch from the endpoint.

Value

A list of class `bco.domain`

Examples

```
fhir_endpoint <- "https://fhirtest.uhn.ca/baseDstu3"
fhir_version <- "3"
fhir_resources <- data.frame(
  "id" = c("21376", "6288583", "25544", "92440", "4588936"),
  "resource" = c(
    "Sequence", "DiagnosticReport", "ProcedureRequest",
    "Observation", "FamilyMemberHistory"
  ),
  stringsAsFactors = FALSE
)

compose_fhir(fhir_endpoint, fhir_version, fhir_resources) %>% convert_json()
```

compose_io_v1.4.2 *Compose BioCompute Object - Input and Output Domain (v1.4.2)*

Description

This domain contains the list of global input and output files created by the computational workflow, excluding the intermediate files.

Usage

```
compose_io_v1.4.2(input = NULL, output = NULL)
```

```
compose_io(input = NULL, output = NULL)
```

Arguments

input	Data frame. Variables include filename, uri, and access_time. Each row is one item in the input subdomain.
output	Data frame. Variables include mediatype, uri, and access_time. Each row is one item in the output subdomain.

Value

A list of class `bco.domain`

Examples

```
input_subdomain <- data.frame(
  "filename" = c(
    "Hepatitis C virus genotype 1",
    "Hepatitis C virus type 1b complete genome"
  ),
  "uri" = c(
    "https://www.ncbi.nlm.nih.gov/nuccore/22129792",
    "https://www.ncbi.nlm.nih.gov/nuccore/5420376"
  ),
  "access_time" = c(
    as.POSIXct("2017-01-24T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST"),
    as.POSIXct("2017-01-24T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST")
  ),
  stringsAsFactors = FALSE
)
```

```
output_subdomain <- data.frame(
  "mediatype" = c("text/csv", "text/csv"),
  "uri" = c(
    "https://example.com/data/514769/dnaAccessionBased.csv",
    "https://example.com/data/514801/SNPPProfile*.csv"
  ),
)
```

```

    "access_time" = c(
      as.POSIXct("2017-01-24T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST"),
      as.POSIXct("2017-01-24T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST")
    ),
    stringsAsFactors = FALSE
  )

  compose_io(input_subdomain, output_subdomain) %>% convert_json()

```

compose_parametric_v1.4.2

Compose BioCompute Object - Parametric Domain (v1.4.2)

Description

Non-default parameters customizing the computational flow which can affect the output of the calculations ([domain definition](#)).

Usage

```
compose_parametric_v1.4.2(df = NULL)
```

```
compose_parametric(df = NULL)
```

Arguments

df Data frame. Variables include param (parameter names), value (value of the parameters), and step (step number for each parameter).

Value

A list of class `bco.domain`

Examples

```

df_parametric <- data.frame(
  "param" = c(
    "seed", "minimum_match_len",
    "divergence_threshold_percent",
    "minimum_coverage", "freq_cutoff"
  ),
  "value" = c("14", "66", "0.30", "15", "0.10"),
  "step" = c(1, 1, 1, 2, 2)
)

compose_parametric(df_parametric) %>% convert_json()

```

compose_provenance_v1.4.2

Compose BioCompute Object - Provenance Domain (v1.4.2)

Description

Compose BioCompute Object - Provenance Domain (v1.4.2)

Usage

```
compose_provenance_v1.4.2(  
  name = NULL,  
  version = NULL,  
  review = NULL,  
  derived_from = NULL,  
  obsolete_after = NULL,  
  embargo = NULL,  
  created = NULL,  
  modified = NULL,  
  contributors = NULL,  
  license = NULL  
)
```

```
compose_provenance(  
  name = NULL,  
  version = NULL,  
  review = NULL,  
  derived_from = NULL,  
  obsolete_after = NULL,  
  embargo = NULL,  
  created = NULL,  
  modified = NULL,  
  contributors = NULL,  
  license = NULL  
)
```

Arguments

name	Character string. Name for the BCO.
version	Character string. Version of this BCO instance object. Should follow the Semantic Versioning format (MAJOR.MINOR.PATCH).
review	Data frame. Reviewer identifiers and descriptions of the status of an object in the review process.
derived_from	Character string. Inheritance/derivation description.
obsolete_after	Date-time object. Expiration date of the object (optional).

embargo	Vector of date-time objects <code>start_time</code> and <code>end_time</code> . If the object has a period of time that it is not public, that range can be specified with this.
created	Date-time object. Initial creation time of the object.
modified	Date-time object. The most recent modification time of the object.
contributors	Data frame. Contributor identifiers and descriptions of their contribution types.
license	Character string. Licence URL or other licence information (text).

Value

A list of class `bco.domain`

Examples

```
name <- "HCV1a ledipasvir resistance SNP detection"
version <- "1.0.0"
review <- data.frame(
  "status" = c("approved", "approved"),
  "reviewer_comment" = c(
    "Approved by [company name] staff. Waiting for approval from FDA Reviewer",
    "The revised BCO looks fine"
  ),
  "date" = c(
    as.POSIXct("2017-11-12T12:30:48", format = "%Y-%m-%dT%H:%M:%S", tz = "EST"),
    as.POSIXct("2017-12-12T12:30:48", format = "%Y-%m-%dT%H:%M:%S", tz = "America/Los_Angeles")
  ),
  "reviewer_name" = c("Jane Doe", "John Doe"),
  "reviewer_affiliation" = c("Seven Bridges Genomics", "U.S. Food and Drug Administration"),
  "reviewer_email" = c("example@sevenbridges.com", "example@fda.gov"),
  "reviewer_contribution" = c("curatedBy", "curatedBy"),
  "reviewer_orcid" = c("https://orcid.org/0000-0000-0000-0000", NA),
  stringsAsFactors = FALSE
)

derived_from <- "https://github.com/biocompute-objects/BCO_Specification/blob/1.2.1-beta/HCV1a.json"
obsolete_after <- as.POSIXct("2018-11-12T12:30:48", format = "%Y-%m-%dT%H:%M:%S", tz = "EST")

embargo <- c(
  "start_time" = as.POSIXct("2017-10-12T12:30:48", format = "%Y-%m-%dT%H:%M:%S", tz = "EST"),
  "end_time" = as.POSIXct("2017-11-12T12:30:48", format = "%Y-%m-%dT%H:%M:%S", tz = "EST")
)

created <- as.POSIXct("2017-01-20T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST")

modified <- as.POSIXct("2019-05-10T09:40:17", format = "%Y-%m-%dT%H:%M:%S", tz = "EST")

contributors <- data.frame(
  "name" = c("Jane Doe", "John Doe"),
  "affiliation" = c("Seven Bridges Genomics", "U.S. Food and Drug Administration"),
  "email" = c("example@sevenbridges.com", "example@fda.gov"),
  "contribution" = I(list(c("createdBy", "curatedBy"), c("authoredBy"))),
  "orcid" = c("https://orcid.org/0000-0000-0000-0000", NA),
```

```

  stringsAsFactors = FALSE
)

license <- "https://creativecommons.org/licenses/by/4.0/"

compose_provenance(
  name, version, review, derived_from, obsolete_after,
  embargo, created, modified, contributors, license
) %>% convert_json()

```

compose_scm_v1.4.2 *Compose BioCompute Object - SCM Extension (v1.4.2)*

Description

Compose BioCompute Object - SCM Extension (v1.4.2)

Usage

```

compose_scm_v1.4.2(
  scm_repository = NULL,
  scm_type = c("git", "svn", "hg", "other"),
  scm_commit = NULL,
  scm_path = NULL,
  scm_preview = NULL
)

compose_scm(
  scm_repository = NULL,
  scm_type = c("git", "svn", "hg", "other"),
  scm_commit = NULL,
  scm_path = NULL,
  scm_preview = NULL
)

```

Arguments

scm_repository	Character string. Base URL of the SCM repository.
scm_type	Character string. Type of SCM database. Must be one of "git", "svn", "hg", or "other".
scm_commit	Character string. Revision within the SCM repository. Should be a repository-wide commit identifier or name of a tag, but may be a name of a branch.
scm_path	Character string. Path from the repository to the source code referenced. Should not start with /.
scm_preview	Character string. The full URI for the source code referenced by the BioCompute Object.

Value

A list of class `bco.domain`

Examples

```
scm_repository <- "https://github.com/example/repo"
scm_type <- "git"
scm_commit <- "c9ffea0b60fa3bcf8e138af7c99ca141a6b8fb21"
scm_path <- "workflow/hive-viral-mutation-detection.cwl"
scm_preview <- "https://github.com/example/repo/blob/master/mutation-detection.cwl"

compose_scm(scm_repository, scm_type, scm_commit, scm_path, scm_preview) %>% convert_json()
```

compose_tlf_v1.4.2 *Compose BioCompute Object - Top Level Fields (v1.4.2)*

Description

Compose BioCompute Object - Top Level Fields (v1.4.2)

Usage

```
compose_tlf_v1.4.2(  
  provenance,  
  usability,  
  extension,  
  description,  
  execution,  
  parametric,  
  io,  
  error,  
  object_id = NULL  
)
```

```
compose_tlf(  
  provenance,  
  usability,  
  extension,  
  description,  
  execution,  
  parametric,  
  io,  
  error,  
  object_id = NULL  
)
```

Arguments

provenance	Provenance domain
usability	Usability domain
extension	Extension domain
description	Description domain
execution	Execution domain
parametric	Parametric domain
io	I/O domain
error	Error domain
object_id	BioCompute Object identifier (definition). If NULL, will use a UUID generated by <code>generate_id</code> .

Value

A vector of top level fields

Examples

```
compose_tlf(
  compose_provenance(), compose_usability(), compose_extension(),
  compose_description(), compose_execution(), compose_parametric(),
  compose_io(), compose_error()
) %>% convert_json()
```

compose_usability_v1.4.2

Compose BioCompute Object - Usability Domain (v1.4.2)

Description

The usability domain (**domain definition**).

Usage

```
compose_usability_v1.4.2(text = NULL)
```

```
compose_usability(text = NULL)
```

Arguments

text	A character vector of free text values that could improves search-ability, provide specific scientific use cases, and a description of the function of the object.
------	--

Value

A list of class `bco.domain`

Examples

```

text <- c(
  paste(
    "Identify baseline single nucleotide polymorphisms (SNPs)[SO:0000694]",
    "(insertions)[SO:0000667], and (deletions)[SO:0000045] that correlate",
    "with reduced (ledipasvir)[pubchem.compound:67505836] antiviral drug",
    "efficacy in (Hepatitis C virus subtype 1)[taxonomy:31646]"
  ),
  paste(
    "Identify treatment emergent amino acid (substitutions)[SO:1000002]",
    "that correlate with antiviral drug treatment failure"
  ),
  paste(
    "Determine whether the treatment emergent amino acid",
    "(substitutions)[SO:1000002] identified correlate with treatment",
    "failure involving other drugs against the same virus"
  )
)

text %>%
  compose_usability() %>%
  convert_json()

```

 compose_v1.4.2

Compose BioCompute Object (v1.4.2)

Description

Compose BioCompute Object (v1.4.2)

Usage

```

compose_v1.4.2(
  tlf,
  provenance,
  usability,
  extension,
  description,
  execution,
  parametric,
  io,
  error
)

compose(
  tlf,
  provenance,
  usability,

```

```

    extension,
    description,
    execution,
    parametric,
    io,
    error
  )

```

Arguments

tlf	Top level fields
provenance	Provenance domain
usability	Usability domain
extension	Extension domain
description	Description domain
execution	Execution domain
parametric	Parametric domain
io	I/O domain
error	Error domain

Value

A list of class bco

Examples

```

tlf <- compose_tlf(
  compose_provenance(), compose_usability(), compose_extension(),
  compose_description(), compose_execution(), compose_parametric(),
  compose_io(), compose_error()
)
biocompute::compose(
  tlf,
  compose_provenance(), compose_usability(), compose_extension(),
  compose_description(), compose_execution(), compose_parametric(),
  compose_io(), compose_error()
) %>% convert_json()

```

convert_json

Convert BioCompute Object or domain to JSON string

Description

Convert BioCompute Object or domain to JSON string

Usage

```
convert_json(x, pretty = TRUE, auto_unbox = TRUE, na = "string", ...)
```

Arguments

x	BioCompute Object or domain
pretty	Prettify the JSON string? Default is TRUE.
auto_unbox	Unbox all atomic vectors of length 1? Default is TRUE.
na	How to represent NA values: must be "null" or "string". Default is "string".
...	Additional parameters for toJSON .

Value

JSON string of the BioCompute Object

Examples

```
compose_description() %>% convert_json()
generate_example("minimal") %>% convert_json()
```

```
convert_yaml
```

Convert BioCompute Object or domain to YAML string

Description

Convert BioCompute Object or domain to YAML string

Usage

```
convert_yaml(x, ...)
```

Arguments

x	BioCompute Object or domain
...	Additional parameters for as.yaml .

Value

YAML string of the BioCompute Object

Examples

```
compose_description() %>%
  convert_yaml() %>%
  cat()
generate_example("minimal") %>%
  convert_yaml() %>%
  cat()
```

export_html	<i>Export BioCompute Object as HTML</i>
-------------	---

Description

Export BioCompute Object as HTML

Usage

```
export_html(x, file, wrap = FALSE, linewidth = 80, ...)
```

Arguments

x	BioCompute Object JSON string from convert_json
file	HTML output file path
wrap	Should the long lines be wrapped?
linewidth	Maximum linewidth when wrap is TRUE.
...	Additional parameters for render .

Value

Path to the output file

Examples

```
## Not run:  
file_html <- tempfile(fileext = ".html")  
generate_example("HCV1a") %>%  
  convert_json() %>%  
  export_html(file_html)  
  
## End(Not run)
```

export_json	<i>Export BioCompute Object as JSON</i>
-------------	---

Description

Export BioCompute Object as JSON

Usage

```
export_json(x, file)
```

Arguments

x BioCompute Object JSON string from [convert_json](#)
 file JSON file path

Value

Path to the output file

Examples

```
file_json <- tempfile(fileext = ".json")
generate_example("HCV1a") %>%
  convert_json() %>%
  export_json(file_json)
cat(paste(readLines(file_json), collapse = "\n"))
```

 export_pdf

Export BioCompute Object as PDF

Description

Export BioCompute Object as PDF

Usage

```
export_pdf(x, file, wrap = FALSE, linewidth = 80, ...)
```

Arguments

x BioCompute Object JSON string from [convert_json](#)
 file PDF output file path
 wrap Should the long lines be wrapped?
 linewidth Maximum linewidth when wrap is TRUE.
 ... Additional parameters for [render](#).

Value

Path to the output file

Examples

```
## Not run:
file_pdf <- tempfile(fileext = ".pdf")
generate_example("HCV1a") %>%
  convert_json() %>%
  export_pdf(file_pdf)

## End(Not run)
```

export_sevenbridges *Export BioCompute Object to Seven Bridges Platforms*

Description

Export BioCompute Object to Seven Bridges Platforms

Usage

```
export_sevenbridges(  
  file,  
  name = NULL,  
  project = NULL,  
  token = NULL,  
  base_url = "https://api.sbgenomics.com/v2/",  
  overwrite = TRUE  
)
```

Arguments

file	Path to the BCO file.
name	Name of the BCO file to create on the platform. Defaults to the name of the input file.
project	Project to upload (export) the BCO file to. Format: "username/project".
token	API auth token for the platform. Generate the token from the platform's Developer Dashboard.
base_url	API base URL. Get the base URL from the platform's Developer Dashboard.
overwrite	If TRUE, will overwrite the existing BCO file with the same name in that project (if any). If FALSE, will not overwrite.

Value

Response of the file upload request

Examples

```
## Not run:  
file_json <- tempfile(fileext = ".json")  
generate_example("HCV1a") %>%  
  convert_json() %>%  
  export_json(file_json)  
  
try(  
  export_sevenbridges(  
    file_json,  
    project = "rosalind_franklin/project_name",  
    token = "your_api_auth_token",
```

```
    base_url = "https://cgc-api.sbgenomics.com/v2/"
  )
)

## End(Not run)
```

export_word

Export BioCompute Object as Word document

Description

Export BioCompute Object as Word document

Usage

```
export_word(x, file, wrap = FALSE, linewidth = 80, ...)
```

Arguments

x	BioCompute Object JSON string from convert_json
file	Word (docx) output file path
wrap	Should the long lines be wrapped?
linewidth	Maximum linewidth when wrap is TRUE.
...	Additional parameters for render .

Value

Path to the output file

Examples

```
## Not run:
file_docx <- tempfile(fileext = ".docx")
generate_example("HCV1a") %>%
  convert_json() %>%
  export_word(file_docx)

## End(Not run)
```

generate_example *Generate example BioCompute Objects*

Description

Generate example BioCompute Objects

Usage

```
generate_example(type = c("minimal", "HCV1a"))
```

Arguments

type Example type. Default is "minimal".

Value

Example BioCompute Object

Examples

```
generate_example("minimal") %>% convert_json()
```

generate_id *Generate ID for the BioCompute Object*

Description

Generate ID for the BioCompute Object

Usage

```
generate_id(platform = c("sevenbridges"))
```

Arguments

platform Platform. Default is "sevenbridges".

Value

BioCompute Object ID

Examples

```
generate_id()
```

is_bco	<i>Is this a BCO object?</i>
--------	------------------------------

Description

Is this a BCO object?

Usage

```
is_bco(x)
```

Arguments

x	any object
---	------------

Value

Logical. TRUE if it is a BCO object, FALSE if not.

Examples

```
generate_example("minimal") %>% is_bco()
```

is_domain	<i>Is this a domain object?</i>
-----------	---------------------------------

Description

Is this a domain object?

Usage

```
is_domain(x)
```

Arguments

x	any object
---	------------

Value

Logical. TRUE if it is a domain object, FALSE if not.

Examples

```
is_domain(compose_description())
```

`read_bco`*Parse Biocompute Object From JSON File to R Object*

Description

Parse Biocompute Object From JSON File to R Object

Usage

```
read_bco(x, ...)
```

Arguments

`x` BioCompute Object .json file
`...` Additional parameters for [fromJSON](#).

Value

A list of class bco

Examples

```
bco <- tempfile(fileext = ".json")
bco <- generate_example("HCV1a") %>%
  convert_json() %>%
  export_json(bco)
bco %>% read_bco()
```

`validate_checksum_v1.4.2`*BioCompute Objects checksum validator (v1.4.2)*

Description

BioCompute Objects checksum validator (v1.4.2)

Usage

```
validate_checksum_v1.4.2(file)
```

```
validate_checksum(file)
```

Arguments

`file` Path to the BCO JSON file

Value

Logical. TRUE if the checksum matched, FALSE if not.

Note

An SHA-256 checksum is **calculated and stored** in the top level fields when a BioCompute Object is created. In reality, due to the delicate differences in how the data in JSON is represented, parsed, and handled in different languages, there could be false positives in the validation results.

Examples

```
bco <- tempfile(fileext = ".json")
generate_example("HCV1a") %>%
  convert_json() %>%
  export_json(bco)
bco %>% validate_checksum()
```

validate_schema_v1.4.2

BioCompute Objects schema validator (v1.4.2)

Description

BioCompute Objects schema validator (v1.4.2)

Usage

```
validate_schema_v1.4.2(file)
```

```
validate_schema(file)
```

Arguments

file Path to the BCO JSON file

Value

None

Note

JSON schema validators for BCO domains and complete BCO based on jsonvalidate. Refer to the [BioCompute Objects Schema](#) for specific JSON schemas.

Examples

```
bco <- tempfile(fileext = ".json")
generate_example("HCV1a") %>%
  convert_json() %>%
  export_json(bco)
bco %>% validate_schema()
```

versions

BioCompute Object specification versions

Description

BioCompute Object specification versions

Usage

```
versions()
```

Value

List of current and all available BioCompute Object specification versions supported by the package.

Examples

```
versions()
```

Index

as.yaml, [19](#)

compose (compose_v1.4.2), [17](#)
compose_description
 (compose_description_v1.4.2), [2](#)
compose_description_v1.4.2, [2](#)
compose_error (compose_error_v1.4.2), [5](#)
compose_error_v1.4.2, [5](#)
compose_execution
 (compose_execution_v1.4.2), [6](#)
compose_execution_v1.4.2, [6](#)
compose_extension
 (compose_extension_v1.4.2), [8](#)
compose_extension_v1.4.2, [8](#)
compose_fhir, [8](#)
compose_fhir (compose_fhir_v1.4.2), [9](#)
compose_fhir_v1.4.2, [9](#)
compose_io (compose_io_v1.4.2), [10](#)
compose_io_v1.4.2, [10](#)
compose_parametric
 (compose_parametric_v1.4.2), [11](#)
compose_parametric_v1.4.2, [11](#)
compose_provenance
 (compose_provenance_v1.4.2), [12](#)
compose_provenance_v1.4.2, [12](#)
compose_scm, [8](#)
compose_scm (compose_scm_v1.4.2), [14](#)
compose_scm_v1.4.2, [14](#)
compose_tlf (compose_tlf_v1.4.2), [15](#)
compose_tlf_v1.4.2, [15](#)
compose_usability
 (compose_usability_v1.4.2), [16](#)
compose_usability_v1.4.2, [16](#)
compose_v1.4.2, [17](#)
convert_json, [18](#), [20](#), [21](#), [23](#)
convert_yaml, [19](#)

export_html, [20](#)
export_json, [20](#)
export_pdf, [21](#)
export_sevenbridges, [22](#)
export_word, [23](#)
fromJSON, [26](#)
generate_example, [24](#)
generate_id, [16](#), [24](#)
is_bco, [25](#)
is_domain, [25](#)
read_bco, [26](#)
render, [20](#), [21](#), [23](#)
toJSON, [19](#)
validate_checksum
 (validate_checksum_v1.4.2), [26](#)
validate_checksum_v1.4.2, [26](#)
validate_schema
 (validate_schema_v1.4.2), [27](#)
validate_schema_v1.4.2, [27](#)
versions, [28](#)