

# Package ‘cbpManager’

July 9, 2025

**Type** Package

**Title** Generate, manage, and edit data and metadata files suitable for  
the import in cBioPortal for Cancer Genomics

**Version** 1.16.0

**Date** 2021-03-16

**Description** This R package provides an R Shiny application that enables the user to generate,  
manage, and edit data and metadata files suitable for the import in cBioPortal for Cancer Ge-  
nomics.

Create cancer studies and edit its metadata. Upload mutation data of a patient that will be con-  
catenated to the data\_mutation\_extended.txt file of the study.

Create and edit clinical patient data, sample data, and timeline data. Create custom time-  
line tracks for patients.

**License** AGPL-3 + file LICENSE

**Encoding** UTF-8

**LazyData** true

**RoxygenNote** 7.2.3

**VignetteBuilder** knitr

**URL** <https://arsenij-ust.github.io/cbpManager/index.html>

**BugReports** <https://github.com/arsenij-ust/cbpManager/issues>

**Depends** shiny, shinydashboard

**Imports** utils, DT, htmltools, vroom, plyr, dplyr, magrittr, jsonlite,  
rapportools, basilisk, reticulate, shinyBS, shinyCSSloaders,  
rintrojs, rlang, markdown

**Suggests** knitr, BiocStyle, rmarkdown, testthat (>= 3.0.0)

**StagedInstall** no

**Collate** 'basilisk.R' 'dynamicTable.R' 'modulesResourceButtons.R'  
'cbpManager-pkg.R' 'functions.R' 'global.R' 'cbpManager.R'  
'shinyAppServer.R' 'shinyAppUI.R'

**biocViews** ImmunoOncology, DataImport, DataRepresentation, GUI,  
ThirdPartyClient, Preprocessing, Visualization

**Config/testthat.edition** 3**git\_url** https://git.bioconductor.org/packages/cbpManager**git\_branch** RELEASE\_3\_21**git\_last\_commit** d49dd22**git\_last\_commit\_date** 2025-04-15**Repository** Bioconductor 3.21**Date/Publication** 2025-07-09**Author** Arsenij Ustjanzew [aut, cre, cph] (ORCID:<<https://orcid.org/0000-0002-1014-4521>>),Federico Marini [aut] (ORCID: <<https://orcid.org/0000-0003-3252-7758>>)**Maintainer** Arsenij Ustjanzew <[arsenij.ustjanzew@gmail.com](mailto:arsenij.ustjanzew@gmail.com)>

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---

`addColumn_Server`      *Server logic of module for adding a column*

---

## Description

Server logic of module for adding a column

## Usage

```
addColumn_Server(input, output, session, data)
```

## Arguments

input	Shiny input
output	Shiny output
session	Shiny session
data	source data as data.frame

## Value

reactive data.frame of modified source data

addColumn\_UI

*UI elements of module for adding a column***Description**

UI elements of module for adding a column

**Usage**

```
addColumn_UI(id, label = "Add column")
```

**Arguments**

<code>id</code>	module id
<code>label</code>	label of the button

**Value**

UI module

addRowRc\_Server

*Server logic of Resource tab module for adding a row***Description**

Server logic of Resource tab module for adding a row

**Usage**

```
addRowRc_Server(
  input,
  output,
  session,
  data,
  patient_ids = NULL,
  sample_ids = NULL,
  resource_ids = NULL,
  resource_type = c("definition", "sample", "patient", "study")
)
```

**Arguments**

input	Shiny input
output	Shiny output
session	Shiny session
data	Source data as data.frame
patient_ids	Reactive vector of existing patient IDs
sample_ids	Reactive data.frame of existing patient IDs and sample IDs
resource_ids	Reactive data.frame of data_resource_definition
resource_type	The type of the resource. Can be "definition", "sample", "patient", "study"

**Value**

reactive data.frame of modified source data

---

addRowRc\_UI

*UI elements of Resource tab module for adding a row*

---

**Description**

UI elements of Resource tab module for adding a row

**Usage**

```
addRowRc_UI(id, label = "Add")
```

**Arguments**

id	module id
label	label of the button

**Value**

UI module

---

addRow_Server	<i>Server logic of module for adding a row</i>
---------------	--

---

**Description**

Server logic of module for adding a row

**Usage**

```
addRow_Server(
  input,
  output,
  session,
  data,
  patient_ids = NULL,
  dates_first_diagnosis = NULL,
  mode = c("timeline", "timepoint")
)
```

**Arguments**

input	Shiny input
output	Shiny output
session	Shiny session
data	source data as data.frame
patient_ids	reactive vector of existing patient IDs
dates_first_diagnosis	data.frame with dates of the first diagnosis and patient IDs
mode	Mode of the timeline data. Controls which columns are displayed.

**Value**

reactive data.frame of modified source data

---

addRow_UI	<i>UI elements of module for adding a row</i>
-----------	---

---

**Description**

UI elements of module for adding a row

**Usage**

```
addRow_UI(id, label = "Add")
```

**Arguments**

id	module id
label	label of the button

**Value**

UI module

---

cBioPortalToDataFrame *Convert the cBioPortal sample- and patient-data file format into a data.frame*

---

**Description**

This function takes a file object (from read.table), removes the # symbol, sets the 5th row as the column names of the data.frame and removes the rows containing the priority, data type and column name. use read.table as follows: read.table(file, sep='\t', colClasses = 'character', comment.char = '')

**Usage**

```
cBioPortalToDataFrame(data)
```

**Arguments**

data	The data.frame of a cBioPortal sample/patient data file
------	---

**Value**

data.frame

**Examples**

```
df <- data.frame(
  V1=c("#attr_1", "#attribute 1", "#STRING", "#1", "ATTRIBUTE_1", "value_1"),
  V2=c("attr_2", "attribute 2", "STRING", "1", "ATTRIBUTE_2", "value_2")
)
cbpManager:::cBioPortalToDataFrame(df)
```

---

**cbpManager***Launch cbpManager*

---

**Description**

Launch the cbpManager Shiny application.

**Usage**

```
cbpManager(studyDir = NULL, logDir = NULL, returnAppObj = FALSE, ...)
```

**Arguments**

studyDir	Path to study folder containing studies of cBioPortal.
logDir	Path where a logfile should be saved. If NULL, logs are not stored
returnAppObj	Logical value, whether to return the app object if set to TRUE. Default behavior: directly runs the app (FALSE)
...	Further parameters that are used by shiny::runApp, e.g. host or port.

**Value**

shiny application object

**Examples**

```
if (interactive()) {  
  cbpManager()  
}
```

---

**cbpManager-pkg***cbpManager*

---

**Description**

‘cbpManager’ is an R package that provides an interactive Shiny-based graphical user interface for...

**Author(s)**

Arsenij Ustjanzew <arsenij.ustjanzew@gmail.com>

---

check\_input\_dates      *Check the input of dates*

---

## Description

Check the input of dates

## Usage

```
check_input_dates(diagnosisDate, startDate = NULL, endDate = NULL)
```

## Arguments

diagnosisDate	date of first diagnosis
startDate	start date of timeline event
endDate	end date of timeline event

## Value

Returns a number indicating the warning

## Examples

```
cbpManager:::check_input_dates(  
  diagnosisDate = "2020-01-01",  
  startDate = "2020-02-01",  
  endDate = "2020-03-01"  
) #returns 0  
cbpManager:::check_input_dates(  
  diagnosisDate = "2020-01-01",  
  startDate = "2019-02-01"  
) #returns 2  
cbpManager:::check_input_dates(  
  diagnosisDate = "2020-01-01",  
  endDate = "2019-02-01"  
) #returns 2  
cbpManager:::check_input_dates(  
  diagnosisDate = "2020-01-01",  
  startDate = "2020-03-01",  
  endDate = "2020-02-01"  
) #returns 1
```

`convertDataFrame`*Convert the data.frame to the appropriate file format for cBioPortal***Description**

Convert the data.frame to the appropriate file format for cBioPortal

**Usage**

```
convertDataFrame(df)
```

**Arguments**

<code>df</code>	data.frame
-----------------	------------

**Value**

Data.frame formated for the cBioPortal file format

**Examples**

```
cbpManager:::convertDataFrame(
  data.frame(
    ATTRIBUTE1=c("attr_1", "attribute 1", "STRING", "value_a1"),
    ATTRIBUTE2=c("attr_2", "attribute 2", "STRING", "value_b1")
  )
)
```

`create_name`*Sanitize names***Description**

This function takes a character string, replaces spaces by underscores and runs make.names.

**Usage**

```
create_name(x, toupper = TRUE)
```

**Arguments**

<code>x</code>	A character string.
<code>toupper</code>	If TRUE, the name wil be upper-case; if FALSE, the name will be lower-case.

**Value**

A sanitized string.

**Examples**

```
cbpManager:::create_name("Study name 1") #returns "STUDY_NAME_1"  
cbpManager:::create_name("FANCY;name", toupper = FALSE) #returns "fancy.name"
```

---

deleteColumn\_Server     *Server logic of module for deleting a column*

---

**Description**

Server logic of module for deleting a column

**Usage**

```
deleteColumn_Server(input, output, session, data, exclude)
```

**Arguments**

input	Shiny input
output	Shiny output
session	Shiny session
data	source data as data.frame
exclude	column names that shoud be excluded from deletion

**Value**

reactive data.frame of modified source data

---

deleteColumn\_UI     *UI elements of module for deleting a column*

---

**Description**

UI elements of module for deleting a column

**Usage**

```
deleteColumn_UI(id, label = "Delete column(s)")
```

**Arguments**

<code>id</code>	module id
<code>label</code>	label of the button

**Value**

UI module

---

`deleteRowRc_Server`     *Server logic of Resource tab module for deleting a row*

---

**Description**

Server logic of Resource tab module for deleting a row

**Usage**

```
deleteRowRc_Server(
  input,
  output,
  session,
  data,
  selected_row,
  mode = "default",
  sample_data = NULL,
  patient_data = NULL,
  study_data = NULL
)
```

**Arguments**

<code>input</code>	Shiny input
<code>output</code>	Shiny output
<code>session</code>	Shiny session
<code>data</code>	Source data as data.frame
<code>selected_row</code>	Index of the selected row from the table
<code>mode</code>	If 'recursive' the resources bind to the resource definition will be deleted.
<code>sample_data</code>	Data of the resource from type 'sample'
<code>patient_data</code>	Data of the resource from type 'patient'
<code>study_data</code>	Data of the resource from type 'study'

**Value**

reactive data.frame of modified source data

---

deleteRowRc_UI	<i>UI elements of module for removing a row</i>
----------------	---

---

**Description**

UI elements of module for removing a row

**Usage**

```
deleteRowRc_UI(id, label = "Delete")
```

**Arguments**

id	Module id
label	Label of the button

**Value**

UI module

---

deleteRow_Server	<i>Server logic of module for removing a row</i>
------------------	--

---

**Description**

Server logic of module for removing a row

**Usage**

```
deleteRow_Server(input, output, session, data, selected_row)
```

**Arguments**

input	Shiny input
output	Shiny output
session	Shiny session
data	source data as data.frame
selected_row	Index of the selected row from the table

**Value**

reactive data.frame of modified source data

---

deleteRow_UI	<i>UI elements of module for removing a row</i>
--------------	---

---

### Description

UI elements of module for removing a row

### Usage

```
deleteRow_UI(id, label = "Delete")
```

### Arguments

id	module id
label	label of the button

### Value

UI module

---

editRowRc_Server	<i>Server logic of Resource tab module for editing a row</i>
------------------	--

---

### Description

Server logic of Resource tab module for editing a row

### Usage

```
editRowRc_Server(  
  input,  
  output,  
  session,  
  data,  
  patient_ids = NULL,  
  sample_ids = NULL,  
  resource_ids = NULL,  
  selected_row = NULL,  
  resource_type = c("definition", "sample", "patient", "study")  
)
```

**Arguments**

input	Shiny input
output	Shiny output
session	Shiny session
data	Source data as data.frame
patient_ids	Reactive vector of existing patient IDs
sample_ids	Reactive data.frame of existing patient IDs and sample IDs
resource_ids	Reactive data.frame of data_resource_definition
selected_row	Index of the selected row
resource_type	The type of the resource. Can be "definition", "sample", "patient", "study"

**Value**

reactive data.frame of modified source data

---

editRowRc\_UI

*UI elements of module for editing a row*

---

**Description**

UI elements of module for editing a row

**Usage**

```
editRowRc_UI(id, label = "Edit")
```

**Arguments**

id	Module id
label	Label of the button

**Value**

UI module

---

editRow_Server	<i>Server logic of module for editing a row</i>
----------------	---

---

## Description

Server logic of module for editing a row

## Usage

```
editRow_Server(  
  input,  
  output,  
  session,  
  data,  
  patient_ids = NULL,  
  dates_first_diagnosis = NULL,  
  selected_row = NULL,  
  mode = c("timeline", "timepoint")  
)
```

## Arguments

input	Shiny input
output	Shiny output
session	Shiny session
data	source data as data.frame
patient_ids	reactive vector of existing patient IDs
dates_first_diagnosis	data.frame with dates of the first diagnosis and patient IDs
selected_row	the index of the selected row
mode	Mode of the timeline data. Controls which columns are displayed.

## Value

reactive data.frame of modified source data

---

editRow\_UI

*UI elements of module for editing a row*

---

### Description

UI elements of module for editing a row

### Usage

```
editRow_UI(id, label = "Edit")
```

### Arguments

id	module id
label	label of the button

### Value

UI module

---

fncols

*Add empty column to a data.frame, if column does not exist in the data.frame*

---

### Description

Add empty column to a data.frame, if column does not exist in the data.frame

### Usage

```
fncols(data, cname)
```

### Arguments

data	data.frame
cname	column name

### Value

data.frame

### Examples

```
cbpManager:::fncols(data.frame(a=c(1,2,3), b=c(4,5,6)), "new")
```

**generateOncotreeUIwidgets***Create shiny UI-widget for specific columns of oncotree entries*

---

**Description**

Create shiny UI-widget for specific columns of oncotree entries

**Usage**

```
generateOncotreeUIwidgets(
  colname,
  mode = c("add", "edit"),
  tab = c("Patient", "Sample")
)
```

**Arguments**

colname	column name
mode	determines the inputId prefix of the UI-widget
tab	"Patient", "Sample" - The used tab; sets the html id prefix of the input

**Value**

A oncotree specific shiny UI-widget

**Examples**

```
oncotree <- jsonlite::fromJSON(system.file("extdata", "oncotree.json", package = "cbpManager"))
cancer_type <- unique(oncotree$mainType[which(!is.na(oncotree$mainType))])
cbpManager:::generateOncotreeUIwidgets("CANCER_TYPE", "add")
```

---

**generateUIwidgets**      *Generate UI input widget*

---

**Description**

Generate UI input widget

**Usage**

```
generateUIwidgets(
  colname,
  mode = c("add", "edit"),
  tab = c("Patient", "Sample", "Mutation"),
  data = NULL,
  selected_row = NULL,
  patientIDs = NULL,
  sampleIDs = NULL
)
```

**Arguments**

colname	A character string - the name of the column, that will be the label of the input
mode	"add" or "edit" - whether to use existing values or not
tab	"Patient", "Sample" - The used tab; sets the html id prefix of the input
data	A data.frame.
selected_row	A number indicating the row number of the selected row in the data.frame.
patientIDs	Vector of patient IDs used for drop down menu of the PATIENT_ID column

**Value**

A sanitized string.

**Examples**

```
cbpManager:::generateUIwidgets(colname = "attribute", mode = "add", tab = "Patient")
```

---

getSampleIDs

*Get Sample IDs associated with Patient IDs from the data\_clinical\_sample.txt file*

---

**Description**

Get Sample IDs associated with Patient IDs from the data\_clinical\_sample.txt file

**Usage**

```
getSampleIDs(file_path, patIDs)
```

**Arguments**

file_path	A character string.
patIDs	A character string.

**Value**

vector with Sample IDs

**Examples**

```
cbpManager:::getSampleIDs(
  system.file("study/testpatient/data_clinical_sample.txt", package = "cbpManager"),
  patIDs = "Testpatient")
```

**importPatientData**      *Import patient data into current study data.frames*

**Description**

Import patient data into current study data.frames

**Usage**

```
importPatientData(
  mode = c("patient", "sample", "mutations", "timelines"),
  file_name,
  file_path,
  patIDs,
  data,
  associatedSampleIDs = NULL
)
```

**Arguments**

mode	Defines the type of imported data
file_name	Filename of source data
file_path	Filepath with filename of source data
patIDs	PATIENT_IDs of patients that should be imported
data	Source data, to be subsetted according to patIDs
associatedSampleIDs	The sample IDs associated to the patIDs

**Value**

data.frame

---

IsDate	<i>Check if input is in the appropriate date format</i>
--------	---

---

**Description**

Check if input is in the appropriate date format

**Usage**

```
IsDate(mydate, date.format = "%Y-%m-%d")
```

**Arguments**

mydate	date
date.format	string describibg the date format

**Value**

boolean

**Examples**

```
cbpManager:::IsDate("2020-02-20")
cbpManager:::IsDate("20.01.2020", date.format = "%d.%m.%Y")
```

---

saveResource_Server	<i>Server logic of module for saving the resource data</i>
---------------------	--

---

**Description**

Server logic of module for saving the resource data

**Usage**

```
saveResource_Server(
  input,
  output,
  session,
  data,
  study_id,
  data_filename,
  meta_filename,
  resource_type = c("SAMPLE", "DEFINITION", "PATIENT", "STUDY")
)
```

**Arguments**

<code>input</code>	Shiny input
<code>output</code>	Shiny output
<code>session</code>	Shiny session
<code>data</code>	Source data as data.frame
<code>study_id</code>	The current study ID
<code>data_filename</code>	File name of the data file
<code>meta_filename</code>	file name of the meta file
<code>resource_type</code>	The type of the resource. Can be "definition", "sample", "patient", "study"

**Value**

boolean value; TRUE if function was used.

`saveResource_UI`

*UI elements of module for saving the resource data*

**Description**

UI elements of module for saving the resource data

**Usage**

```
saveResource_UI(id, label = "Save")
```

**Arguments**

<code>id</code>	module id
<code>label</code>	label of the button

**Value**

UI module

---

saveTimeline\_Server     *Server logic of module for saving the source data*

---

### Description

Server logic of module for saving the source data

### Usage

```
saveTimeline_Server(input, output, session, data, study_id)
```

### Arguments

input	Shiny input
output	Shiny output
session	Shiny session
data	source data as data.frame
study_id	the current study ID

### Value

boolean value; TRUE if function was used.

---

saveTimeline\_UI     *UI elements of module for saving the data*

---

### Description

UI elements of module for saving the data

### Usage

```
saveTimeline_UI(id, label = "Save")
```

### Arguments

id	module id
label	label of the button

### Value

UI module

`setupConda_cbpManager` *Install conda environment with basilisk before launching the app*

### Description

Install conda environment with basilisk before launching the app

### Usage

```
setupConda_cbpManager()
```

### Value

Nothing to return

### Examples

```
## Not run:  
setupConda_cbpManager()  
  
## End(Not run)
```

`shinyAppServer` *Shiny app server function*

### Description

Shiny app server function

### Usage

```
shinyAppServer(input, output, session)
```

### Arguments

<code>input</code>	provided by shiny
<code>output</code>	provided by shiny
<code>session</code>	provided by shiny

### Value

nothing to return

---

`shinyAppUI`

*Shiny app server object create the shiny application user interface*

---

### Description

Shiny app server object create the shiny application user interface

### Usage

```
shinyAppUI
```

### Format

An object of class `shiny.tag` of length 3.

---

`updateOncotreeUIwidgets`

*Updates UI-widgets for specific columns of oncotree entries*

---

### Description

Updates UI-widgets for specific columns of oncotree entries

### Usage

```
updateOncotreeUIwidgets(  
  session,  
  row_last_clicked,  
  mode = c("add", "edit"),  
  tab = c("Patient", "Sample")  
)
```

### Arguments

<code>session</code>	Shiny session
<code>row_last_clicked</code>	the index of the row last clicked in the oncotree_table
<code>mode</code>	determines the inputId prefix of the UI-widget
<code>tab</code>	"Patient", "Sample" - The used tab; sets the html id pr

### Value

nothing to return

---

`validateResourceDefinition`

*Validate resource\_definition input*

---

**Description**

Validate resource\_definition input

**Usage**

```
validateResourceDefinition(values, resourceDf, mode = "add")
```

**Arguments**

values	List of input values
resourceDf	data.frame of data_resource_definition
mode	The mode of the function ('add' or 'edit')

**Value**

boolean

---

---

`validateResourcePatient`

*Validate resource\_patient input*

---

**Description**

Validate resource\_patient input

**Usage**

```
validateResourcePatient(values)
```

**Arguments**

values	List of input values
--------	----------------------

**Value**

boolean

---

```
validateResourceSample
```

*Validate resource\_sample input*

---

**Description**

Validate resource\_sample input

**Usage**

```
validateResourceSample(values)
```

**Arguments**

values	List of input values
--------	----------------------

**Value**

boolean
---------

---

```
validateResourceStudy  Validate resource_study input
```

---

**Description**

Validate resource\_study input

**Usage**

```
validateResourceStudy(values)
```

**Arguments**

values	List of input values
--------	----------------------

**Value**

boolean
---------

**ValidationDependencies***Validation Dependencies***Description**

Vector defining a set of Python dependencies and versions required to operate with the validation scripts for cBioPortal

**Usage**

```
.validation_dependencies
```

**Format**

A character vector containing the pinned versions of all Python packages on which the import validation depends.

**writeLogFile**

*Write a line to a logfile containing the date, time, username (from Shinyproxy), and the name of the modified file.*

---

**Description**

Write a line to a logfile containing the date, time, username (from Shinyproxy), and the name of the modified file.

**Usage**

```
writeLogFile(outdir, modified_file, log_filename = "cbpManager_logfile.txt")
```

**Arguments**

outdir            directory, where the logfile should be saved  
modified\_file    Name of the modified file  
log\_filename    Name of the logfile

**Value**

Nothing to return

**Examples**

```
cbpManager:::writeLogFile(tempdir(), "data_clinical_patient.txt")
```

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