Package 'AnVILPublish'

March 29, 2021

Title Publish Packages and Other Resources to AnVIL Workspaces
Version 1.0.0
Description Use this package to create or update AnVIL workspaces from resources such as R / Bioconductor packages. The metadata about the package (e.g., select information from the package DESCRIPTION file and from vignette YAML headings) are used to populate the 'DASHBOARD'. Vignettes are translated to python notebooks ready for evaluation in AnVIL.
License Artistic-2.0
Imports AnVIL, httr, rmarkdown, whisker, tools, utils, stats,
Suggests knitr, BiocStyle, BiocManager
biocViews Infrastructure, Software
Encoding UTF-8
Roxygen list(markdown = TRUE)
RoxygenNote 7.1.1
VignetteBuilder knitr
git_url https://git.bioconductor.org/packages/AnVILPublish
git_branch RELEASE_3_12
git_last_commit fe3b8cb
git_last_commit_date 2020-10-27
Date/Publication 2021-03-29
Author Martin Morgan [aut, cre] (https://orcid.org/0000-0002-5874-8148)
Maintainer Martin Morgan <mtmorgan.bioc@gmail.com></mtmorgan.bioc@gmail.com>
R topics documented:
add_access
Index

2 as_notebook

add_access

Add Bioconductor_User group to workspace access

Description

add_access() adds the Bioconductor_User group to a workspace with READER permissions. Users gain access to the workspace (and others) by being added to the Bioconductor_User group.

Usage

```
add_access(namespace, name)
```

Arguments

namespace

character(1) namespace (billing account) under which the workspace belongs.

name

character(1) name of the workspace to add access credentials.

Value

add_access() returns TRUE, invisibly.

as_notebook

Render vignettes as .ipynb notebooks

Description

as_notebook() renders .Rmd vignettes as .ipynb notebooks, and updates the notebooks in an AnVIL workspace.

Usage

```
as_notebook(rmd_paths, namespace, name, update = FALSE)
```

Arguments

rmd_paths character() paths to to Rmd files.

namespace character(1) AnVIL namespace (billing project) to be used.

name character(1) AnVIL workspace name.

update logical(1) Update (over-write any similarly named notebooks) an existing

workspace? The default (FALSE) creates notebooks locally, e.g., for previewing

via jupyter notebook *ipynb.

Details

.Rmd Vignettes are processed to .md using rmarkdown::render(...,md_document()), and then translated to .ipynb using python software called notedown; notedown is available at https://github.com/aaren/notedown

The translation is not perfect, for instance code chunks marked as `eval = FALSE` are not marked as such in the python notebook.

as_workspace 3

Value

as_notebook() returns the paths to the local (if update = FALSE) or the workspace notebooks.

as_workspace	Render R packages as AnVIL workspaces

Description

as_workspace() renders a package source tree (e.g., from a git checkout) as an AnVIL workspace.

Usage

```
as_workspace(path, namespace, name = NULL, create = FALSE, update = FALSE)
```

Arguments

path character(1) path to the location of the package source code.

namespace character(1) AnVIL namespace (billing project) to be used.

name character(1) AnVIL workspace name or NULL. If NULL, the workspace name is set to "Bioconductor-Package-<pkgname>", where <pkgname> is the name of the package (from the DESCRIPTION file) at path.

create logical(1) Create a new project?

update logical(1) Update (over-write the existing DASHBOARD and any similarly named notebooks) an existing workspace? If niether create nore update is TRUE, the code to create a workspace is run but no output generated; this can be useful during debugging.

Details

Information from the DESCRIPTION file and Rmd YAML files are used to populate the 'DASH-BOARD' tab.

See `?as_notebook()` for details on how vignettes are processed to notebooks, and the limitations of the current approach.

Value

as_workspace() returns the URL of the updated workspace, invisibly.

Index

add_access, 2
as_notebook, 2
as_workspace, 3