Package 'startR'

December 21, 2023

Title Automatically Retrieve Multidimensional Distributed Data Sets

Version 2.3.1

Description Tool to automatically fetch, transform and arrange subsets of multi- dimensional data sets (collections of files) stored in local and/or remote file systems or servers, using multicore capabilities where possible. The tool provides an interface to perceive a collection of data sets as a single large multidimensional data array, and enables the user to request for automatic retrieval, processing and arrangement of subsets of the large array. Wrapper functions to add support for custom file formats can be plugged in/out, making the tool suitable for any research field where large multidimensional data sets are involved.

Depends R (>= 3.6.0)

Imports abind, bigmemory, future, multiApply (>= 2.1.0), parallel, easyNCDF, s2dv, ClimProjDiags, PCICt, methods

Suggests stats, utils, testthat, yaml

License GPL-3

URL https://earth.bsc.es/gitlab/es/startR/

BugReports https://earth.bsc.es/gitlab/es/startR/-/issues

SystemRequirements cdo ecFlow

Encoding UTF-8

RoxygenNote 7.2.3

Config/testthat/edition 3

NeedsCompilation no

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Repository CRAN

Date/Publication 2023-12-21 16:20:02 UTC

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AddStep

Create the workflow with the previous defined operation and data.

Description

The step that combines the previous declared data and operation together to create the complete workflow. It is the final step before data processing.

Usage

```
AddStep(inputs, step_fun, ...)
```

inputs	One or a list of objects of the class 'startR_cube' returned by Start(), indicating the data to be processed.
step_fun	A startR step function as returned by Step().
	Additional parameters for the inputs of function defined in 'step_fun' by Step().

CDORemapper

Value

A list of the class 'startR_workflow' containing all the objects needed for the data operation.

Examples

```
data_path <- system.file('extdata', package = 'startR')</pre>
path_obs <- file.path(data_path, 'obs/monthly_mean/$var$/$var$_$sdate$.nc')</pre>
sdates <- c('200011', '200012')</pre>
data <- Start(dat = list(list(path = path_obs)),</pre>
               var = 'tos',
               sdate = sdates,
               time = 'all',
               latitude = 'all',
              longitude = 'all',
              return_vars = list(latitude = 'dat',
                                   longitude = 'dat',
                                   time = 'sdate'),
               retrieve = FALSE)
pi_short <- 3.14
fun <- function(x, pi_val) {</pre>
          lat = attributes(x)$Variables$dat1$latitude
          weight = sqrt(cos(lat * pi_val / 180))
          corrected = Apply(list(x), target_dims = "latitude",
                              fun = function(x) {x * weight})
        }
step <- Step(fun = fun,</pre>
              target_dims = 'latitude',
              output_dims = 'latitude',
              use_libraries = c('multiApply'),
              use_attributes = list(data = "Variables"))
wf <- AddStep(data, step, pi_val = pi_short)</pre>
```

CDORemapper

CDO Remap Data Transformation for 'startR'

Description

This is a transform function that uses CDO software to remap longitude-latitude data subsets onto a specified target grid, intended for use as parameter 'transform' in a Start() call. This function complies with the input/output interface required by Start() defined in the documentation for the parameter 'transform' of function Start().

This function uses the function CDORemap() in the package 's2dv' to perform the interpolation, hence CDO is required to be installed.

Usage

```
CDORemapper(
   data_array,
   variables,
   file_selectors = NULL,
   crop_domain = NULL,
   ...
)
```

Arguments

data_array	A data array to be transformed. See details in the documentation of the parameter 'transform' of the function Start().
variables	A list of auxiliary variables required for the transformation, automatically provided by Start(). See details in the documentation of the parameter 'transform' of the function Start().
file_selectors	A charcter vector indicating the information of the path of the file parameter 'data_array' comes from. See details in the documentation of the parameter 'transform' of the function Start(). The default value is NULL.
crop_domain	A list of the transformed domain of each transform variable, automatically provided by Start().
	A list of additional parameters to adjust the transform process, as provided in the parameter 'transform_params' in a Start() call. See details in the documentation of the parameter 'transform' of the function Start().

Value

An array with the same amount of dimensions as the input data array, potentially with different sizes, and potentially with the attribute 'variables' with additional auxiliary data. See details in the documentation of the parameter 'transform' of the function Start().

See Also

CDORemap

Examples

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Collect

End(Not run)

Collect

Collect and merge the computation results

Description

The final step of the startR workflow after the data operation. It is used when the parameter 'wait' of Compute() is FALSE. It combines all the chunks of the results as one data array when the execution is done. See more details on practical guide. Collect() calls Collect_ecflow() or Collect_autosubmit() according to the chosen workflow manager.

Usage

```
Collect(startr_exec, wait = TRUE, remove = TRUE, on_remote = FALSE)
```

Arguments

startr_exec	An R object returned by Compute() when the parameter 'wait' of Compute() is FALSE. It can be directly from a Compute() call or read from the RDS file.
wait	A logical value deciding whether the R session waits for the Collect() call to finish (TRUE) or not (FALSE). If TRUE, it will be a blocking call, in which Collect() will retrieve information from the HPC, including signals and outputs, each polling_period seconds. The the status can be monitored on the workflow manager GUI. Collect() will not return until the results of all the chunks have been received. If FALSE, Collect() return an error if the execution has not finished, otherwise it will return the merged array. The default value is TRUE.
remove	A logical value deciding whether to remove of all chunk results received from the HPC after data being collected, as well as the local job folder under 'ecflow_suite_dir or 'autosubmit_suite_dir'. To preserve the data and Collect() them as many times as desired, set remove to FALSE. The default value is TRUE.
on_remote	A logical value deciding to the function is run locally and sync the outputs back from HPC (FALSE, default), or it is run on HPC (TRUE).

Value

A list of merged data array.

Examples

```
data_path <- system.file('extdata', package = 'startR')</pre>
path_obs <- file.path(data_path, 'obs/monthly_mean/$var$/$var$_$sdate$.nc')</pre>
sdates <- c('200011', '200012')</pre>
data <- Start(dat = list(list(path = path_obs)),</pre>
              var = 'tos',
              sdate = sdates,
              time = 'all',
               latitude = 'all',
              longitude = 'all',
               return_vars = list(latitude = 'dat',
                                   longitude = 'dat',
                                   time = 'sdate'),
              retrieve = FALSE)
fun <- function(x) {
          lat = attributes(x)$Variables$dat1$latitude
          weight = sqrt(cos(lat * pi / 180))
          corrected = Apply(list(x), target_dims = "latitude",
                             fun = function(x) {x * weight})
        }
step <- Step(fun = fun,</pre>
              target_dims = 'latitude',
             output_dims = 'latitude',
             use_libraries = c('multiApply'),
             use_attributes = list(data = "Variables"))
wf <- AddStep(data, step)</pre>
## Not run:
res <- Compute(wf, chunks = list(longitude = 2, sdate = 2),</pre>
                threads_load = 1,
                threads_compute = 4,
                cluster = list(queue_host = 'nord3',
                                queue_type = 'lsf',
                                temp_dir = '/on_hpc/tmp_dir/',
                                cores_per_job = 2,
                                job_wallclock = '05:00',
                                max_jobs = 4,
                                extra_queue_params = list('#BSUB -q bsc_es'),
                                bidirectional = FALSE,
                               polling_period = 10
                ),
                ecflow_suite_dir = '/on_local_machine/username/ecflow_dir/',
                wait = FALSE)
saveRDS(res, file = 'test_collect.Rds')
collect_info <- readRDS('test_collect.Rds')</pre>
result <- Collect(collect_info, wait = TRUE)</pre>
```

End(Not run)

Compute

Specify the execution parameters and trigger the execution

Compute

Description

The step of the startR workflow after the complete workflow is defined by AddStep(). This function specifies the execution parameters and triggers the execution. The execution can be operated locally or on a remote machine. If it is the latter case, the configuration of the machine needs to be sepecified in the function, and the EC-Flow server is required to be installed.

The execution can be operated by chunks to avoid overloading the RAM memory. After all the chunks are finished, Compute() will gather and merge them, and return a single data object, including one or multiple multidimensional data arrays and additional metadata.

Usage

```
Compute(
  workflow,
  chunks = "auto",
  workflow_manager = "ecFlow",
  threads_load = 1,
  threads_compute = 1,
  cluster = NULL,
  ecflow_suite_dir = NULL,
  ecflow_server = NULL,
  autosubmit_suite_dir = NULL,
  autosubmit_server = NULL,
  silent = FALSE,
  debug = FALSE,
  wait = TRUE
)
```

workflow	A list of the class 'startR_workflow' returned by function AddSteop() or of class 'startR_cube' returned by function Start(). It contains all the objects needed for the execution.	
chunks	A named list of dimensions which to split the data along and the number of chunks to make for each. The chunked dimension can only be those not required as the target dimension in function Step(). The default value is 'auto', which lists all the non-target dimensions and each one has one chunk.	
workflow_manager		
	Can be NULL, 'ecFlow' or 'Autosubmit'. The default is 'ecFlow'.	
threads_load	An integer indicating the number of parallel execution cores to use for the data retrieval stage. The default value is 1.	
threads_compute		
	An integer indicating the number of parallel execution cores to use for the computation. The default value is 1.	
cluster	A list of components that define the configuration of the machine to be run on. The comoponents vary from the different machines. Check Practical guide on GitLab for more details and examples. Only needed when the computation is not run locally. The default value is NULL.	

ecflow_suite_dir		
	A character string indicating the path to a folder in the local workstation where to store temporary files generated for the automatic management of the workflow. Only needed when the execution is run remotely. The default value is NULL.	
ecflow_server	A named vector indicating the host and port of the EC-Flow server. The vector form should be c(host = 'hostname', port = port_number). Only needed when the execution is run remotely. The default value is NULL.	
autosubmit_suit	e_dir	
	A character string indicating the path to a folder where to store temporary files generated for the automatic management of the workflow manager. This path should be available in local workstation as well as autosubmit machine. The default value is NULL, and a temporary folder under the current working folder will be created.	
autosubmit_serv	ver in the second s	
	A character vector indicating the login node of the autosubmit machine. It can be "bscesautosubmit01" or "bscesautosubmit02". The default value is NULL, and the node will be randomly chosen.	
silent	A logical value deciding whether to print the computation progress (FALSE) on the R session or not (TRUE). It only works when the execution runs locally or the parameter 'wait' is TRUE. The default value is FALSE.	
debug	A logical value deciding whether to return detailed messages on the progress and operations in a Compute() call (TRUE) or not (FALSE). Automatically changed to FALSE if parameter 'silent' is TRUE. The default value is FALSE.	
wait	A logical value deciding whether the R session waits for the Compute() call to finish (TRUE) or not (FALSE). If FALSE, it will return an object with all the information of the startR execution that can be stored in your disk. After that, the R session can be closed and the results can be collected later with the Collect() function. The default value is TRUE.	

Value

A list of data arrays for the output returned by the last step in the specified workflow (wait = TRUE), or an object with information about the startR execution (wait = FALSE). The configuration details and profiling information are attached as attributes to the returned list of arrays.

indices

indices

Specify dimension selectors with indices

Description

This is a helper function used in a Start() call to define the desired range of dimensions. It selects the indices of the coordinate variable from original data. See details in the documentation of the parameter . . . 'indices to take' of the function Start().

Usage

indices(x)

Arguments

Х

A numeric vector or a list with two nemerics to take all the elements between the two specified indices (both extremes inclusive).

Value

Same as input, but with additional attribute 'indices', 'values', and 'chunk'.

See Also

values

```
# Used in Start():
    data_path <- system.file('extdata', package = 'startR')
    path_obs <- file.path(data_path, 'obs/monthly_mean/$var$/$var$_$sdate$.nc')
    sdates <- c('200011', '200012')
    data <- Start(dat = list(list(path = path_obs)),</pre>
```

NcCloser

NetCDF file closer for 'startR'

Description

This is a file closer function for NetCDF files, intended for use as parameter 'file_closer' in a Start() call. This function complies with the input/output interface required by Start() defined in the documentation for the parameter 'file_closer'.

This function uses the function NcClose() in the package 'easyNCDF', which in turn uses nc_close() in the package 'ncdf4'.

Usage

NcCloser(file_object)

Arguments

file_object An open connection to a NetCDF file, optionally with additional header information. See details in the documentation of the parameter 'file_closer' of the function Start().

Value

This function returns NULL.

See Also

NcOpener NcDataReader NcDimReader NcVarReader

```
data_path <- system.file('extdata', package = 'startR')
path_obs <- file.path(data_path, 'obs/monthly_mean/tos/tos_200011.nc')
connection <- NcOpener(path_obs)
NcCloser(connection)</pre>
```

NcDataReader

Description

This is a data reader function for NetCDF files, intended for use as parameter file_data_reader in a Start() call. This function complies with the input/output interface required by Start() defined in the documentation for the parameter 'file_data_reader'.

This function uses the function NcToArray() in the package 'easyNCDF', which in turn uses nc_var_get() in the package 'ncdf4'.

Usage

```
NcDataReader(
   file_path = NULL,
   file_object = NULL,
   file_selectors = NULL,
   inner_indices = NULL,
   synonims
)
```

Arguments

file_path	A character string indicating the path to the data file to read. See details in the documentation of the parameter 'file_data_reader' of the function Start(). The default value is NULL.
file_object	An open connection to a NetCDF file, optionally with additional header infor- mation. See details in the documentation of the parameter 'file_data_reader' of the function Start(). The default value is NULL.
file_selectors	A named list containing the information of the path of the file to read data from. It is automatically provided by Start(). See details in the documentation of the parameter 'file_data_reader' of the function Start(). The default value is NULL.
inner_indices	A named list of numeric vectors indicating the indices to take from each of the inner dimensions in the requested file. It is automatically provided by Start(). See details in the documentation of the parameter 'file_data_reader' of the function Start(). The default value is NULL.
synonims	A named list indicating the synonims for the dimension names to look for in the requested file, exactly as provided in the parameter 'synonims' in a Start() call. See details in the documentation of the parameter 'file_data_reader' of the function Start().

Value

A multidimensional data array with the named dimensions and indices requested in 'inner_indices', potentially with the attribute 'variables' with additional auxiliary data. See details in the documentation of the parameter 'file_data_reader' of the function Start().

See Also

NcOpener NcDimReader NcCloser NcVarReader

Examples

```
NcDimReader
```

NetCDF dimension reader for 'startR'

Description

A dimension reader function for NetCDF files, intended for use as parameter 'file_dim_reader' in a Start() call. It complies with the input/output interface required by Start() defined in the documentation for the parameter 'file_dim_reader' of that function.

This function uses the function NcReadDims() in the package 'easyNCDF'.

Usage

```
NcDimReader(
   file_path = NULL,
   file_object = NULL,
   file_selectors = NULL,
   inner_indices = NULL,
   synonims
)
```

file_path	A character string indicating the path to the data file to read. See details in the documentation of the parameter 'file_dim_reader' of the function Start(). The default value is NULL.
file_object	An open connection to a NetCDF file, optionally with additional header infor- mation. See details in the documentation of the parameter 'file_dim_reader' of the function Start(). The default value is NULL.
file_selectors	A named list containing the information of the path of the file to read data from. It is automatically provided by Start(). See details in the documentation of the parameter 'file_dim_reader' of the function Start(). The default value is NULL.

NcOpener

inner_indices	A named list of numeric vectors indicating the indices to take from each of the inner dimensions in the requested file. It is automatically provided by Start(). See details in the documentation of the parameter 'file_dim_reader' of the function Start(). The default value is NULL.
synonims	A named list indicating the synonims for the dimension names to look for in the requested file, exactly as provided in the parameter 'synonims' in a Start() call. See details in the documentation of the parameter 'file_dim_reader' of the function Start().

Value

A named numeric vector with the names and sizes of the dimensions of the requested file.

See Also

NcOpener NcDataReader NcCloser NcVarReader

Examples

NcOpener

NetCDF file opener for 'startR'

Description

This is a file opener function for NetCDF files, intended for use as parameter 'file_opener' in a Start() call. This function complies with the input/output interface required by Start() defined in the documentation for the parameter 'file_opener'.

This function uses the function NcOpen() in the package 'easyNCDF', which in turn uses nc_open() in the package 'ncdf4'.

Usage

```
NcOpener(file_path)
```

file_path	A character string indicating the path to the data file to read. See details in the
	documentation of the parameter 'file_opener' of the function Start().

Value

An open connection to a NetCDF file with additional header information as returned by nc_open() in the package 'ncdf4'. See details in the documentation of the parameter 'file_opener' of the function Start().

See Also

NcDimReader NcDataReader NcCloser NcVarReader

Examples

```
data_path <- system.file('extdata', package = 'startR')
path_obs <- file.path(data_path, 'obs/monthly_mean/tos/tos_200011.nc')
connection <- NcOpener(path_obs)
NcCloser(connection)</pre>
```

NcVarReader

NetCDF variable reader for 'startR'

Description

This is an auxiliary variable reader function for NetCDF files, intended for use as parameter 'file_var_reader' in a Start() call. It complies with the input/output interface required by Start() defined in the documentation for the parameter 'file_var_reader' of that function.

This function uses the function NcDataReader() in the package 'startR', which in turn uses Nc-ToArray() in the package 'easyNCDF', which in turn uses nc_var_get() in the package 'ncdf4'.

Usage

```
NcVarReader(
   file_path = NULL,
   file_object = NULL,
   file_selectors = NULL,
   var_name = NULL,
   synonims
)
```

file_path	A character string indicating the path to the data file to read the variable from. See details in the documentation of the parameter 'file_var_reader' of the function Start(). The default value is NULL.
file_object	An open connection to a NetCDF file, optionally with additional header infor- mation. See details in the documentation of the parameter 'file_var_reader' of the function Start(). The default value is NULL.

file_selectors	A named list containing the information of the path of the file to read data from. It is automatically provided by Start(). See details in the documentation of the parameter 'file_var_reader' of the function Start(). The default value is NULL.
var_name	A character string with the name of the variable to be read. The default value is NULL.
synonims	A named list indicating the synonims for the dimension names to look for in the requested file, exactly as provided in the parameter 'synonims' in a Start() call. See details in the documentation of the parameter 'file_var_reader' of the function Start().

Value

A multidimensional data array with the named dimensions, potentially with the attribute 'variables' with additional auxiliary data. See details in the documentation of the parameter 'file_var_reader' of the function Start().

See Also

NcOpener NcDataReader NcCloser NcDimReader

Examples

SelectorChecker Translate a set of selectors into a set of numeric indices

Description

This is a selector checker function intended for use as parameter 'selector_checker' in a Start() call. It translates a set of selectors which is the value for one dimension into a set of numeric indices corresponding to the coordinate variable. The function complies with the input/output interface required by Start() defined in the documentation for the parameter 'selector_checker' of Start().

Usage

```
SelectorChecker(selectors, var = NULL, return_indices = TRUE, tolerance = NULL)
```

Arguments

selectors	A vector or a list of two of numeric indices or variable values to be retrieved for a dimension, automatically provided by Start(). See details in the documentation of the parameters 'selector_checker' and '' of the function Start().
var	A vector of values of a coordinate variable for which to search matches with the provided indices or values in the parameter 'selectors', automatically provided by Start(). See details in the documentation of the parameters 'selector_checker' and '' of the function Start(). The default value is NULL. When not specified, SelectorChecker() simply returns the input indices.
return_indices	A logical value automatically configured by Start(), telling whether to return the numeric indices or coordinate variable values after the matching. The default value is TRUE.
tolerance	A numeric value indicating a tolerance value to be used in the matching of 'se- lectors' and 'var'. See documentation on ' <dim_name>_tolerance' in in the documentation of the function Start(). The default value is NULL.</dim_name>

Value

A vector of either the indices of the matching values (if return_indices = TRUE) or the matching values themselves (if return_indices = FALSE).

Examples

```
# Get the latitudes from 10 to 20 degree
sub_array_of_selectors <- list(10, 20)
# The latitude values from original file
sub_array_of_values <- seq(90, -90, length.out = 258)[2:257]
SelectorChecker(sub_array_of_selectors, sub_array_of_values)
```

Sort

Sort the coordinate variable values in a Start() call

Description

The reorder function intended for use as parameter '<dim_name>_reorder' in a call to the function Start(). This function complies with the input/output interface required by Start() defined in the documentation for the parameter ... of that function.

The coordinate applied to Sort() consists of an increasing or decreasing sort of the values. It is useful for adjusting the latitude order.

The coordinate applied to CircularSort() consists of a circular sort of values, where any values beyond the limits specified in the parameters 'start' and 'end' is applied a modulus to fall in the specified range. This is useful for circular coordinates such as the Earth longitudes.

Start

Usage

```
Sort(...)
```

```
CircularSort(start, end, ...)
```

Arguments

	Additional parameters to adjust the reorderig. See function sort() for more de- tails.
start	A numeric indicating the lower bound of the circular range.
end	A numeric indicating the upper bound of the circular range.

Value

A list of 2 containing:

\$x	The reordered values.
\$ix	The permutation indices of \$x in the original coordinate.

Examples

```
# Used in Start():
data_path <- system.file('extdata', package = 'startR')</pre>
path_obs <- file.path(data_path, 'obs/monthly_mean/$var$/$var$_$sdate$.nc')</pre>
sdates <- c('200011', '200012')</pre>
data <- Start(dat = list(list(path = path_obs)),</pre>
               var = 'tos',
               sdate = sdates,
               time = 'all',
               latitude = values(list(-60, 60)),
               latitude_reorder = Sort(decreasing = TRUE),
               longitude = values(list(-120, 120)),
               longitude_reorder = CircularSort(-180, 180),
               return_vars = list(latitude = 'dat',
                                    longitude = 'dat',
                                    time = 'sdate'),
               retrieve = FALSE)
```

Start

Declare, discover, subset and retrieve multidimensional distributed data sets

Description

See the startR documentation and tutorial for a step-by-step explanation on how to use Start().

Nowadays in the era of big data, large multidimensional data sets from diverse sources need to be combined and processed. Analysis of big data in any field is often highly complex and time-consuming. Taking subsets of these data sets and processing them efficiently become an indispensable practice. This technique is also known as Domain Decomposition, Map Reduce or, more commonly, 'chunking'.

startR (Subset, TrAnsform, ReTrieve, arrange and process large multidimensional data sets in R) is an R project started at BSC with the aim to develop a tool that allows the user to automatically process large multidimensional distributed data sets. It is an open source project that is open to external collaboration and funding, and will continuously evolve to support as many data set formats as possible while maximizing its efficiency.

startR provides a framework under which a data set (collection of one or multiple data files, potentially distributed over various remote servers) are perceived as if they all were part of a single large multidimensional array. Once such multidimensional array is declared, any user-defined function can be applied to the data in a apply-like fashion, where startR transparently implements the Map Reduce paradigm. The steps to follow in order to process a collection of big data sets are as follows:

- Declaring the data set, i.e. declaring the distribution of the data files involved, the dimensions and shape of the multidimensional array, and the boundaries of the target data. This step can be performed with the Start() function. Numeric indices or coordinate values can be used when fixing the boundaries. It is common having the need to apply transformations, pre-processing or reordering to the data. Start() accepts user-defined transformation or reordering functions to be applied for such purposes. Once a data set is declared, a list of involved files, dimension lengths, memory size and other metadata is made available. Optionally, the data set can be retrieved and loaded onto the current R session if it is small enough.
- Declaring the workflow of operations to perform on the involved data set(s). This step can be performed with the Step() and AddStep() functions.
- Defining the computation settings. The mandatory settings include a) how many subsets to divide the data sets into and along which dimensions; b) which platform to perform the work-flow of operations on (local machine or remote machine/HPC?), how to communicate with it (unidirectional or bidirectional connection? shared or separate file systems?), which queuing system it uses (slurm, PBS, LSF, none?); and c) how many parallel jobs and execution threads per job to use when running the calculations. This step can be performed when building up the call to the Compute() function.
- Running the computation. startR transparently implements the Map Reduce paradigm, according to the settings in the previous steps. The progress can optionally be monitored with the EC-Flow workflow management tool. When the computation ends, a report of performance timings is displayed. This step can be triggered with the Compute() function.

startR is not bound to a specific file format. Interface functions to custom file formats can be provided for Start() to read them. As this version, startR includes interface functions to the following file formats:

Start

• NetCDF

Metadata and auxilliary data is also preserved and arranged by Start() in the measure that it is retrieved by the interface functions for a specific file format.

Usage

```
Start(
  . . . ,
  return_vars = NULL,
  synonims = NULL,
  file_opener = NcOpener,
  file_var_reader = NcVarReader,
  file_dim_reader = NcDimReader,
  file_data_reader = NcDataReader,
  file_closer = NcCloser.
  transform = NULL,
  transform_params = NULL,
  transform_vars = NULL,
  transform_extra_cells = 2,
  apply_indices_after_transform = FALSE,
  pattern_dims = NULL,
 metadata_dims = NULL,
  selector_checker = SelectorChecker,
 merge_across_dims = FALSE,
 merge_across_dims_narm = TRUE,
  split_multiselected_dims = FALSE,
  path_glob_permissive = FALSE,
  largest_dims_length = FALSE,
  retrieve = FALSE,
  num_procs = 1,
 ObjectBigmemory = NULL,
  silent = FALSE,
  debug = FALSE
```

Arguments

)

. . .

A selection of custemized parameters depending on the data format. When we retrieve data from one or a collection of data sets, the involved data can be perceived as belonging to a large multi-dimensional array. For instance, let us consider an example case. We want to retrieve data from a source, which contains data for the number of monthly sales of various items, and also for their retail price each month. The data on source is stored as follows:

#/data/ # |-> sales/ # | |-> electronics

```
# | | |-> item_a.data
# | | |-> item_b.data
# | | |-> item_c.data
# | |-> clothing
# | |-> item_d.data
# | |-> idem_e.data
# | |-> idem_f.data
# |-> prices/
# |-> electronics
# | |-> item_a.data
# | |-> item_b.data
# | |-> item_c.data
# |-> clothing
# |-> item_d.data
# |-> item_e.data
# |-> item_f.data
```

Each item file contains data, stored in whichever format, for the sales or prices over a time period, e.g. for the past 24 months, registered at 100 different stores over the world. Whichever the format it is stored in, each file can be perceived as a container of a data array of 2 dimensions, time and store. Let us assume the '.data' format allows to keep a name for each of these dimensions, and the actual names are 'time' and 'store'.

The different item files for sales or prices can be perceived as belonging to an 'item' dimension of length 3, and the two groups of three items to a 'section' dimension of length 2, and the two groups of two sections (one with the sales and the other with the prices) can be perceived as belonging also to another dimension 'variable' of length 2. Even the source can be perceived as belonging to a dimension 'source' of length 1.

All in all, in this example, the whole data could be perceived as belonging to a multidimensional 'large array' of dimensions

```
# source variable section item store month
# 1 2 2 3 100 24
```

The dimensions of this 'large array' can be classified in two types. The ones that group actual files (the file dimensions) and the ones that group data values inside the files (the inner dimensions). In the example, the file dimensions are 'source', 'variable', 'section' and 'item', whereas the inner dimensions are 'store' and 'month'.

Having the dimensions of our target sources in mind, the parameter . . . expects to receive information on:

- The names of the expected dimensions of the 'large dataset' we want to retrieve data from
- The indices to take from each dimension (and other constraints)

- How to reorder the dimension if needed
- · The location and organization of the files of the data sets

For each dimension, the 3 first information items can be specified with a set of parameters to be provided through For a given dimension 'dimname', six parameters can be specified:

dimname = <indices_to_take>, # 'all' / 'first' / 'last' /
indices(c(1, 10, 20)) /
indices(c(1:20)) /
indices(list(1, 20)) /
c(1, 10, 20) / c(1:20) /
list(1, 20)
dimname_var = <name_of_associated_coordinate_variable>,
dimname_tolerance = <tolerance_value>,
dimname_reorder = <reorder_function>,
dimname_depends = <name_of_another_dimension>,
dimname_across = <name_of_another_dimension>

The indices to take can be specified in three possible formats (see code comments above for examples). The first format consists in using character tags, such as 'all' (take all the indices available for that dimension), 'first' (take only the first) and 'last' (only the last). The second format consists in using numeric indices, which have to be wrapped in a call to the indices() helper function. For the second format, either a vector of numeric indices can be provided, or a list with two numeric indices can be provided to take all the indices in the range between the two specified indices (both extremes inclusive). The third format consists in providing a vector character strings (for file dimensions) or of values of whichever type (for inner dimensions). For the file dimensions, the provided character strings in the third format will be used as components to build up the final path to the files (read further). For inner dimensions, the provided values in the third format will be compared to the values of an associated coordinate variable (must be specified in '<dimname> reorder', read further), and the indices of the closest values will be retrieved. When using the third format, a list with two values can also be provided to take all the indices of the values within the specified range.

The **name of the associated coordinate variable** must be a character string with the name of an associated coordinate variable to be found in the data files (in all* of them). For this to work, a 'file_var_reader' function must be specified when calling Start() (see parameter 'file_var_reader'). The coordinate variable must also be requested in the parameter 'return_vars' (see its section for details). This feature only works for inner dimensions.

The **tolerance value** is useful when indices for an inner dimension are specified in the third format (values of whichever type). In that case, the indices of the closest values in the coordinate variable are seeked. However the closest value might be too distant and we would want to consider no real match exists for such provided value. This is possible via the tolerance, which allows to specify a threshold beyond which not to seek for matching values and mark that index as missing value.

The **reorder_function** is useful when indices for an inner dimension are specified in the third fromat, and the retrieved indices need to be reordered in function of their provided associated variable values. A function can be provided, which receives as input a vector of values, and returns as outputs a list with the components \$x with the reordered values, and \$ix with the permutation indices. Two reordering functions are included in startR, the Sort() and the CircularSort().

The **name of another dimension** to be specified in <dimname>_depends, only available for file dimensions, must be a character string with the name of another requested **file dimension** in ..., and will make Start() aware that the path components of a file dimension can vary in function of the path component of another file dimension. For instance, in the example above, specifying item_depends = 'section' will make Start() aware that the item names vary in function of the section, i.e. section 'electronics' has items 'a', 'b' and 'c' but section 'clothing' has items 'd', 'e', 'f'. Otherwise Start() would expect to find the same item names in all the sections. If values() is used to define dimensions, it is possible to provide different values of the depending dimension for each depended dimension values. For example, if section = c('electronics', 'clothing'), we can use item = list(electronics = c('a', 'b', 'c'), clothing = c('d', 'e', 'f')).

The **name of another dimension** to be specified in '<dimname>_across', only available for inner dimensions, must be a character string with the name of another requested **inner dimension** in ..., and will make Start() aware that an inner dimension extends along multiple files. For instance, let us imagine that in the example above, the records for each item are so large that it becomes necessary to split them in multiple files each one containing the registers for a different period of time, e.g. in 10 files with 100 months each ('item_a_period1.data', 'item_a_period2.data', and so on). In that case, the data can be perceived as having an extra file dimension, the 'period' dimension. The inner dimension 'month' would extend across multiple files, and providing the parameter month = indices(1, 300) would make Start() crash because it would perceive we have made a request out of bounds (each file contains 100 'month' indices, but we requested 1 to 300). This can be solved by specifying the parameter month_across = period (a long with the full specification of the dimension 'period').

Defining the path pattern

As mentioned above, the parameter ... also expects to receive information with the location of the data files. In order to do this, a special dimension must be defined. In that special dimension, in place of specifying indices to take, a path pattern must be provided. The path pattern is a character string that encodes the way the files are organized in their source. It must be a path to one of the data set files in an accessible local or remote file system, or a URL to one of the files provided by a local or remote server. The regions of this path that vary across files (along the file dimensions) must be replaced by wildcards. The wildcards must match any of the defined file dimensions in the call to Start() and must be delimited with heading and trailing '\$'. Shell globbing expressions can be used in the path pattern. See the next code snippet for an example of a path pattern.

All in all, the call to Start() to load the entire data set in the example of store item sales, would look as follows:

```
# data <- Start(source = paste0('/data/$variable$/',
# '$section$/$item$.data'),
# variable = 'all',
# section = 'all',
# item_depends = 'section',
# store = 'all',
# month = 'all')</pre>
```

Note that in this example it would still be pending to properly define the parameters 'file_opener', 'file_closer', 'file_dim_reader', 'file_var_reader' and 'file_data_reader' for the '.data' file format (see the corresponding sections).

The call to Start() will return a multidimensional R array with the following dimensions:

```
# source variable section item store month
# 1 2 2 3 100 24
The dimension specifications in the ... do not have to follow any particular or-
der. The returned array will have the dimensions in the same order as they have
been specified in the call. For example, the following call:
```

```
# data <- Start(source = paste0('/data/$variable$/',
# '$section$/$item$.data'),
# month = 'all',
# store = 'all',
# item_depends = 'section',
# section = 'all',
# variable = 'all')</pre>
```

would return an array with the following dimensions:

```
# source month store item section variable
# 1 24 100 3 2 2
```

Next, a more advanced example to retrieve data for only the sales records, for the first section ('electronics'), for the 1st and 3rd items and for the stores located in Barcelona (assuming the files contain the variable 'store_location' with the name of the city each of the 100 stores are located at):

```
# data <- Start(source = paste0('/data/$variable$/',
# '$section$/$item$.data'),
# variable = 'sales',
# section = 'first',
# item = indices(c(1, 3)),
# item_depends = 'section',
# store = 'Barcelona',
# store_var = 'store_location',
# month = 'all',
# return_vars = list(store_location = NULL))
```

The defined names for the dimensions do not necessarily have to match the names of the dimensions inside the file. Lists of alternative names to be seeked can be defined in the parameter 'synonims'.

If data from multiple sources (not necessarily following the same structure) has to be retrieved, it can be done by providing a vector of character strings with path pattern specifications, or, in the extended form, by providing a list of lists with the components 'name' and 'path', and the name of the dataset and path pattern as values, respectively. For example:

```
# data <- Start(source = list(</pre>
#list(name = 'sourceA',
# path = paste0('/sourceA/$variable$/',
# '$section$/$item$.data')),
#list(name = 'sourceB',
# path = paste0('/sourceB/$section$/',
# '$variable$/$item$.data'))
#),
# variable = 'sales',
# section = 'first',
# item = indices(c(1, 3)),
# item_depends = 'section',
# store = 'Barcelona',
# store_var = 'store_location',
# month = 'all',
# return_vars = list(store_location = NULL))
```

```
return_vars A named list where the names are the names of the variables to be fetched in the files, and the values are vectors of character strings with the names of the file dimension which to retrieve each variable for, or NULL if the variable has to be retrieved only once from any (the first) of the involved files.
```

Apart from retrieving a multidimensional data array, retrieving auxiliary variables inside the files can also be needed. The parameter 'return_vars' allows for requesting such variables, as long as a 'file_var_reader' function is also specified in the call to Start() (see documentation on the corresponding parameter). In the case of the the item sales example (see documentation on parameter \ldots), the store location variable is requested with the parameter

return_vars = list(store_location = NULL).

This will cause Start() to fetch once the variable 'store_location' and return it in the component

\$Variables\$common\$store_location,

and will be an array of character strings with the location names, with the dimensions c('store' = 100). Although useless in this example, we could ask Start() to fetch and return such variable for each file along the items dimension as follows:

return_vars = list(store_location = c('item')).

In that case, the variable will be fetched once from a file of each of the items, and will be returned as an array with the dimensions c('item' = 3, 'store' = 100).

If a variable is requested along a file dimension that contains path pattern specifications ('source' in the example), the fetched variable values will be returned in the component

\$Variables\$<dataset_name>\$<variable_name>.
For example:

```
# data <- Start(source = list(</pre>
#list(name = 'sourceA',
# path = paste0('/sourceA/$variable$/',
# '$section$/$item$.data')),
#list(name = 'sourceB',
# path = paste0('/sourceB/$section$/',
# '$variable$/$item$.data'))
#),
# variable = 'sales',
# section = 'first',
# item = indices(c(1, 3)),
# item_depends = 'section',
# store = 'Barcelona',
# store_var = 'store_location',
# month = 'all',
# return_vars = list(store_location = c('source',
# 'item')))
# # Checking the structure of the returned variables
# str(found_data$Variables)
# Named list
# .. $common: NULL
# ..$sourceA: Named list
# ....$store_location: char[1:18(3d)] 'Barcelona' 'Barcelona' ...
# ..$sourceB: Named list
# ....$store_location: char[1:18(3d)] 'Barcelona' 'Barcelona' ...
# # Checking the dimensions of the returned variable
```

	<pre># # for the source A # dim(found_data\$Variables\$sourceA) # item store # 3 3</pre>
	The names of the requested variables do not necessarily have to match the actual variable names inside the files. A list of alternative names to be seeked can be specified via the parameter 'synonims'.
synonims	A named list where the names are the requested variable or dimension names, and the values are vectors of character strings with alternative names to seek for such dimension or variable.
	In some requests, data from different sources may follow different naming con- ventions for the dimensions or variables, or even files in the same source could have varying names. This parameter is in order for Start() to properly identify the dimensions or variables with different names.
	In the example used in parameter 'return_vars', it may be the case that the two involved data sources follow slightly different naming conventions. For exam- ple, source A uses 'sect' as name for the sections dimension, whereas source B uses 'section'; source A uses 'store_loc' as variable name for the store locations, whereas source B uses 'store_location'. This can be taken into account as fol- lows:
	<pre># data <- Start(source = list(# list(name = 'sourceA', # path = paste@('/sourceA/\$variable\$/', # '\$section\$/\$item\$.data')), # list(name = 'sourceB', # path = paste@('/sourceB/\$section\$/', # '\$variable\$/\$item\$.data')) #), # variable = 'sales', # section = 'first', # item = indices(c(1, 3)), # item_depends = 'section', # store = 'Barcelona', # store_var = 'store_location', # month = 'all', # return_vars = list(store_location = c('source', # 'item')), # synonims = list(# section = c('sec', 'section'), # store_location = c('store_loc', # 'store_location') #))</pre>

file_opener A function that receives as a single parameter 'file_path' a character string with

the path to a file to be opened, and returns an object with an open connection to the file (optionally with header information) on success, or returns NULL on failure.

This parameter takes by default NcOpener() (an opener function for NetCDF files).

See NcOpener() for a template to build a file opener for your own file format.

file_var_reader

A function with the header file_path = NULL, file_object = NULL, file_selectors = NULL, var_name, synonims that returns an array with auxiliary data (i.e. data from a variable) inside a file. Start() will provide automatically either a 'file path' or a 'file object' to the 'file var reader' function (the function has to be ready to work whichever of these two is provided). The parameter 'file selectors' will also be provided automatically to the variable reader, containing a named list where the names are the names of the file dimensions of the queried data set (see documentation on ...) and the values are single character strings with the components used to build the path to the file being read (the one provided in 'file_path' or 'file_object'). The parameter 'var_name' will be filled in automatically by Start() also, with the name of one of the variales to be read. The parameter 'synonims' will be filled in with exactly the same value as provided in the parameter 'synonims' in the call to Start(), and has to be used in the code of the variable reader to check for alternative variable names inside the target file. The 'file var reader' must return a (multi)dimensional array with named dimensions, and optionally with the attribute 'variales' with other additional metadata on the retrieved variable.

Usually, the 'file_var_reader' should be a degenerate case of the 'file_data_reader' (see documentation on the corresponding parameter), so it is recommended to code the 'file_data_reder' in first place.

This parameter takes by default NcVarReader() (a variable reader function for NetCDF files).

See NcVarReader() for a template to build a variale reader for your own file format.

file_dim_reader

A function with the header file_path = NULL, file_object = NULL, file_selectors = NULL, synonims that returns a named numeric vector where the names are the names of the dimensions of the multidimensional data array in the file and the values are the sizes of such dimensions. Start() will provide automatically either a 'file_path' or a 'file_object' to the 'file_dim_reader' function (the function has to be ready to work whichever of these two is provided). The parameter 'file_selectors' will also be provided automatically to the dimension reader, containing a named list where the names are the names of the file dimensions of the queried data set (see documentation on ...) and the values are single character strings with the components used to build the path to the file being read (the one provided in 'file_path' or 'file_object'). The parameter 'synonims' will be filled in with exactly the same value as provided in the parameter 'synonims' in the call to Start(), and can optionally be used in advanced configurations.

This parameter takes by default NcDimReader() (a dimension reader function for NetCDF files).

See NcDimReader() for (an advanced) template to build a dimension reader for your own file format.

file_data_reader

A function with the header file_path = NULL, file_object = NULL, file_selectors = NULL, inner_indices = NULL, synonims that returns a subset of the multidimensional data array inside a file (even if internally it is not an array). Start() will provide automatically either a 'file_path' or a 'file_object' to the 'file_data_reader' function (the function has to be ready to work whichever of these two is provided). The parameter 'file_selectors' will also be provided automatically to the data reader, containing a named list where the names are the names of the file dimensions of the queried data set (see documentation on ...) and the values are single character strings with the components used to build the path to the file being read (the one provided in 'file_path' or 'file_object'). The parameter 'inner indices' will be filled in automatically by Start() also, with a named list of numeric vectors, where the names are the names of all the expected inner dimensions in a file to be read, and the numeric vectors are the indices to be taken from the corresponding dimension (the indices may not be consecutive nor in order). The parameter 'synonims' will be filled in with exactly the same value as provided in the parameter 'synonims' in the call to Start(), and has to be used in the code of the data reader to check for alternative dimension names inside the target file. The 'file_data_reader' must return a (multi)dimensional array with named dimensions, and optionally with the attribute 'variables' with other additional metadata on the retrieved data.

Usually, 'file_data_reader' should use 'file_dim_reader' (see documentation on the corresponding parameter), so it is recommended to code 'file_dim_reder' in first place.

This parameter takes by default NcDataReader() (a data reader function for NetCDF files).

See NcDataReader() for a template to build a data reader for your own file format.

file_closer A function that receives as a single parameter 'file_object' an open connection (as returned by 'file_opener') to one of the files to be read, optionally with header information, and closes the open connection. Always returns NULL.

This parameter takes by default NcCloser() (a closer function for NetCDF files).

See NcCloser() for a template to build a file closer for your own file format.

transform A function with the header dara_array, variables, file_selectors = NULL, It receives as input, through the parameter data_array, a subset of a mul-

tidimensional array (as returned by 'file_data_reader'), applies a transformation to it and returns it, preserving the amount of dimensions but potentially modifying their size. This transformation may require data from other auxiliary variables, automatically provided to 'transform' through the parameter 'variables', in the form of a named list where the names are the variable names and the values are (multi)dimensional arrays. Which variables need to be sent to 'transform' can be specified with the parameter 'transform_vars' in Start(). The parameter 'file_selectors' will also be provided automatically to 'transform', containing a named list where the names are the names of the file dimensions of the queried data set (see documentation on . . .) and the values are single character strings with the components used to build the path to the file the subset being processed belongs to. The parameter . . . will be filled in with other additional parameters to adjust the transformation, exactly as provided in the call to Start() via the parameter 'transform_params'.

transform_params

A named list with additional parameters to be sent to the 'transform' function (if specified). See documentation on parameter 'transform' for details.

transform_vars A vector of character strings with the names of auxiliary variables to be sent to the 'transform' function (if specified). All the variables to be sent to 'transform' must also have been requested as return variables in the parameter 'return_vars' of Start().

transform_extra_cells

An integer of extra indices to retrieve from the data set, beyond the requested indices in ..., in order for 'transform' to dispose of additional information to properly apply whichever transformation (if needed). As many as 'transform_extra_cells' will be retrieved beyond each of the limits for each of those inner dimensions associated to a coordinate variable and sent to 'transform' (i.e. present in 'transform_vars'). After 'transform' has finished, Start() will take again and return a subset of the result, for the returned data to fall within the specified bounds in The default value is 2.

apply_indices_after_transform

A logical value indicating when a 'transform' is specified in Start() and numeric indices are provided for any of the inner dimensions that depend on coordinate variables, these numeric indices can be made effective (retrieved) before applying the transformation or after. The boolean flag allows to adjust this behaviour. It takes FALSE by default (numeric indices are applied before sending data to 'transform').

- pattern_dims A character string indicating the name of the dimension with path pattern specifications (see ... for details). If not specified, Start() assumes the first provided dimension is the pattern dimension, with a warning.
- metadata_dims A vector of character strings with the names of the file dimensions which to return metadata for. As noted in 'file_data_reader', the data reader can optionally return auxiliary data via the attribute 'variables' of the returned array. Start() by default returns the auxiliary data read for only the first file of each source (or data set) in the pattern dimension (see . . . for info on what the pattern dimension is). However it can be configured to return the metadata for all the files along any set of file dimensions. The default value is NULL, and it will be assigned automatically as parameter 'pattern_dims'.

A function used internaly by Start() to translate a set of selectors (values for a dimension associated to a coordinate variable) into a set of numeric indices. It takes by default SelectorChecker() and, in principle, it should not be required to change it for customized file formats. The option to replace it is left open for more versatility. See the code of SelectorChecker() for details on the inputs, functioning and outputs of a selector checker.

merge_across_dims

A logical value indicating whether to merge dimensions across which another dimension extends (according to the '<dimname>_across' parameters). Takes the value FALSE by default. For example, if the dimension 'time' extends across the dimension 'chunk' and merge_across_dims = TRUE, the resulting data array will only contain only the dimension 'time' as long as all the chunks together.

merge_across_dims_narm

A logical value indicating whether to remove the additional NAs from data when parameter 'merge_across_dims' is TRUE. It is helpful when the length of the tobe-merged dimension is different across another dimension. For example, if the dimension 'time' extends across dimension 'chunk', and the time length along the first chunk is 2 while along the second chunk is 10. Setting this parameter as TRUE can remove the additional 8 NAs at position 3 to 10. The default value is TRUE, but will be automatically turned to FALSE if 'merge_across_dims = FALSE'.

split_multiselected_dims

A logical value indicating whether to split a dimension that has been selected with a multidimensional array of selectors into as many dimensions as present in the selector array. The default value is FALSE.

path_glob_permissive

A logical value or an integer specifying how many folder levels in the path pattern, beginning from the end, the shell glob expressions must be preserved and worked out for each file. The default value is FALSE, which is equivalent to 0. TRUE is equivalent to 1.

When specifying a path pattern for a dataset, it might contain shell glob experissions. For each dataset, the first file matching the path pattern is found, and the found file is used to work out fixed values for the glob expressions that will be used for all the files of the dataset. However, in some cases, the values of the shell glob expressions may not be constant for all files in a dataset, and they need to be worked out for each file involved.

For example, a path pattern could be as follows:

'/path/to/dataset/\$var\$_*/\$date\$_*_foo.nc'.

Leaving path_glob_permissive = FALSE will trigger automatic seek of the contents to replace the asterisks (e.g. the first asterisk matches with 'bar' and the second with 'baz'. The found contents will be used for all files in the dataset (in the example, the path pattern will be fixed to

'/path/to/dataset/\$var\$_bar/\$date\$_baz_foo.nc'. However, if any of the files in the dataset have other contents in the position of the asterisks, Start() will not find them (in the example, a file like

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'/path/to/dataset/precipitation_bar/19901101_bin_foo.nc' would not be found). Setting path_glob_permissive = 1 would preserve global expressions in the latest level (in the example, the fixed path pattern would be '/path/to/dataset/\$var\$_bar/\$date\$_*_foo.nc', and the problematic file mentioned before would be found), but of course this would slow down the Start() call if the dataset involves a large number of files. Setting path_glob_permissive = 2 would leave the original path pattern with the original glob expressions in the 1st and 2nd levels (in the example, both asterisks would be preserved, thus would allow Start() to recognize files such as

'/path/to/dataset/precipitation_zzz/19901101_yyy_foo.nc').

Note that each glob expression can only represent one possibility (Start() chooses the first). Because * is not the tag, which means it cannot be a dimension of the output array. Therefore, only one possibility can be adopted. For example, if '/path/to/dataset/precipitation_*/19901101_*_foo.nc' has two matches:

'/path/to/dataset/precipitation_xxx/19901101_yyy_foo.nc' and '/path/to/dataset/precipitation_zzz/19901101_yyy_foo.nc', only the first found file will be used.

largest_dims_length

A logical value or a named integer vector indicating if Start() should examine all the files to get the largest length of the inner dimensions (TRUE) or use the first valid file of each dataset as the returned dimension length (FALSE). Since examining all the files could be time-consuming, a vector can be used to explicitly specify the expected length of the inner dimensions. For those inner dimensions not specified, the first valid file will be used. The default value is FALSE.

This parameter is useful when the required files don't have consistent inner dimension. For example, there are 10 required experimental data files of a series of start dates. The data only contain 25 members for the first 2 years while 51 members for the later years. If 'largest_dims_length = FALSE', the returned member dimension length will be 25 only. The 26th to 51st members in the later 8 years will be discarded. If 'largest_dims_length = TRUE', the returned member dimension length will be 51. To save the resource, 'largest_dims_length = c(member = 51)' can also be used.

- retrieve A logical value indicating whether to retrieve the data defined in the Start() call or to explore only its dimension lengths and names, and the values for the file and inner dimensions. The default value is FALSE.
- num_procs An integer of number of processes to be created for the parallel execution of the retrieval/transformation/arrangement of the multiple involved files in a call to Start(). If set to NULL, takes the number of available cores (as detected by future::availableCores). The default value is 1 (no parallel execution).

ObjectBigmemory

a character string to be included as part of the bigmemory object name. This parameter is thought to be used internally by the chunking capabilities of startR.

silent A logical value of whether to display progress messages (FALSE) or not (TRUE). The default value is FALSE. debug A logical value of whether to return detailed messages on the progress and operations in a Start() call (TRUE) or not (FALSE). The default value is FALSE.

Value

If retrieve = TRUE the involved data is loaded into RAM memory and an object of the class 'startR_cube' with the following components is returned:

Data	Multidimensional data array with named dimensions, with the data values re- quested via and other parameters. This array can potentially contain meta- data in the attribute 'variables'.
Variables	Named list of 1 + N components, containing lists of retrieved variables (as re- quested in 'return_vars') common to all the data sources (in the 1st component, \$common), and for each of the N dara sources (named after the source name, as specified in, or, if not specified, \$dat1, \$dat2,, \$datN). Each of the vari- ables are contained in a multidimensional array with named dimensions, and potentially with the attribute 'variables' with additional auxiliary data.
Files	Multidimensional character string array with named dimensions. Its dimensions are the file dimensions (as requested in). Each cell in this array contains a path to a retrieved file, or NULL if the corresponding file was not found.
NotFoundFiles	Array with the same shape as \$Files but with NULL in the positions for which the corresponding file was found, and a path to the expected file in the positions for which the corresponding file was not found.
FileSelectors	Multidimensional character string array with named dimensions, with the same shape as \$Files and \$NotFoundFiles, which contains the components used to build up the paths to each of the files in the data sources.

If retrieve = FALSE the involved data is not loaded into RAM memory and an object of the class 'startR_header' with the following components is returned:

- Dimensions Named vector with the dimension lengths and names of the data involved in the Start() call.
- Variables Named list of 1 + N components, containing lists of retrieved variables (as requested in 'return_vars') common to all the data sources (in the 1st component, \$common), and for each of the N dara sources (named after the source name, as specified in ..., or, if not specified, \$dat1, \$dat2, ..., \$datN). Each of the variables are contained in a multidimensional array with named dimensions, and potentially with the attribute 'variables' with additional auxiliary data.
- Files Multidimensional character string array with named dimensions. Its dimensions are the file dimensions (as requested in ...). Each cell in this array contains a path to a file to be retrieved (which may exist or not).
- FileSelectors Multidimensional character string array with named dimensions, with the same shape as \$Files and \$NotFoundFiles, which contains the components used to build up the paths to each of the files in the data sources.
- StartRCall List of parameters sent to the Start() call, with the parameter 'retrieve' set to TRUE. Intended for calling in order to retrieve the associated data a posteriori with a call to do.call().

Step

Examples

Step

Define the operation applied on declared data.

Description

The step of the startR workflow after declaring data by Start() call. It identifies the operation (i.e., function) and the target and output dimensions of data array for the function. Ideally, it expects the dimension name to be in the same order as the one requested in the Start() call. If a different order is specified, startR will reorder the subset dimension to the expected order for this function.

Usage

```
Step(
  fun,
  target_dims,
  output_dims,
  use_libraries = NULL,
  use_attributes = NULL
)
```

fun	A function in R format defining the operation to be applied to the data declared by a Start() call. It should only work on the essential dimensions rather than all the data dimensions. Since the function will be called numerous times through all the non-essential dimensions, it is recommended to keep them as light as possible.
target_dims	A vector for single input array or a list of vectors for multiple input arrays indi- cating the names of the dimensions 'fun' to be applied along.
output_dims	A vector for single returned array or a list of vectors for multiple returned arrays indicating the dimension names of the function output.

use_libraries	A vector of character string indicating the R library names to be used in 'fun'. Only used when the jobs are run on HPCs; if the jobs are run locally, load the necessary libraries by library() directly. The default value is NULL.
use_attributes	One or more lists of vectors of character string indicating the data attributes to be used in 'fun'. The list name should be consistent with the list name of 'data' in AddStep(). The default value is NULL.

Value

A closure that contains all the objects assigned. It serves as the input of Addstep().

Examples

```
data_path <- system.file('extdata', package = 'startR')</pre>
path_obs <- file.path(data_path, 'obs/monthly_mean/$var$/$var$_$sdate$.nc')</pre>
sdates <- c('200011', '200012')</pre>
data <- Start(dat = list(list(path = path_obs)),</pre>
              var = 'tos',
               sdate = sdates,
               time = 'all',
               latitude = 'all',
               longitude = 'all',
               return_vars = list(latitude = 'dat',
                                   longitude = 'dat',
                                   time = 'sdate'),
               retrieve = FALSE)
fun <- function(x) {</pre>
          lat = attributes(x)$Variables$dat1$latitude
          weight = sqrt(cos(lat * pi / 180))
          corrected = Apply(list(x), target_dims = "latitude",
                              fun = function(x) {x * weight})
        }
step <- Step(fun = fun,</pre>
              target_dims = 'latitude',
              output_dims = 'latitude',
             use_libraries = c('multiApply'),
             use_attributes = list(data = "Variables"))
wf <- AddStep(data, step)</pre>
```

values

Specify dimension selectors with actual values

Description

This is a helper function used in a Start() call to define the desired range of dimensions. It specifies the actual value to be matched with the coordinate variable. See details in the documentation of the parameter ... 'indices to take' of the function Start().

values

Usage

values(x)

Arguments

х

A numeric vector or a list with two nemerics to take all the element between the two specified values (both extremes inclusive).

Value

Same as input, but with additional attribute 'indices', 'values', and 'chunk'.

See Also

indices

```
# Used in Start():
data_path <- system.file('extdata', package = 'startR')</pre>
path_obs <- file.path(data_path, 'obs/monthly_mean/$var$/$var$_$sdate$.nc')</pre>
sdates <- c('200011', '200012')</pre>
data <- Start(dat = list(list(path = path_obs)),</pre>
               var = 'tos',
               sdate = sdates,
               time = 'all',
               latitude = values(seq(-80, 80, 20)),
               latitude_reorder = Sort(),
               longitude = values(list(10, 300)),
               longitude_reorder = CircularSort(0, 360),
               return_vars = list(latitude = 'dat',
                                   longitude = 'dat',
                                    time = 'sdate'),
                retrieve = FALSE)
```

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