Package 'squarebrackets'

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Type Package

Title Subset Methods as Alternatives to the Square Brackets Operators for Programming

Version 0.0.0.9

Description Provides subset methods

(supporting both atomic and recursive S3 classes)

that may be more convenient alternatives to the `[` and `[<-` operators, whilst maintaining similar performance.

Some nice properties of these methods include, but are not limited to, the following.

1) The `[` and `[<-` operators use different rule-sets for different data.frame-like types (data.frames, data.tables, tibbles, tidytables, etc.).

The 'squarebrackets' methods use the same rule-sets for the different data.frame-like types.

2) Performing dimensional subset operations on an array using `[` and `[<-`, requires a-priori knowledge on the number of dimensions the array has.

The 'squarebrackets' methods work on any arbitrary dimensions without requiring such prior knowledge.

3) When selecting names with the `[` and `[<-` operators,

only the first occurrence of the names are selected in case of duplicate names.

The 'squarebrackets' methods always perform on all names in case of duplicates, not just the first.

4) The `[[` and `[[<-` operators

allow operating on a recursive subset of a nested list.

But these only operate on a single recursive subset,

and are not vectorized for multiple recursive subsets of a nested list at once.

'squarebrackets' provides a way to reshape a nested list

into a recursive matrix,

thereby allowing vectorized operations on recursive subsets of such a nested list.

5) The `[<-` operator only supports copy-on-modify semantics for most classes.

The 'squarebrackets' methods provides explicit pass-by-reference and pass-by-value semantics, whilst still respecting things like binding-locks and mutability rules.

6) 'squarebrackets' supports index-less sub-set operations,

which is more memory efficient

(and better for the environment)

for `long vectors` than sub-set operations using the `[` and `[<-` operators.

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aaa00_squarebrackets_help

squarebrackets: Subset Methods as Alternatives to the Square Brackets Operators for Programming

Description

squarebrackets:

Subset Methods as Alternatives to the Square Brackets Operators for Programming.

'squarebrackets' provides subset methods (supporting both atomic and recursive S3 classes) that may be more convenient alternatives to the [and [<- operators, whilst maintaining similar performance.

Some nice properties of these methods include, but are not limited to, the following.

- 1. The [and [<- operators use different rule-sets for different data.frame-like types (data.frames, data.tables, tibbles, tidytables, etc.).
 - The 'squarebrackets' methods use the same rule-sets for the different data.frame-like types.
- 2. Performing dimensional subset operations on an array using [and [<-, requires a-priori knowledge on the number of dimensions the array has.
 - The 'squarebrackets' methods work on any arbitrary dimensions without requiring such prior knowledge.
- 3. When selecting names with the [and [<- operators, only the first occurrence of the names are selected in case of duplicate names.
 - The 'squarebrackets' methods always perform on all names in case of duplicates, not just the first.
- 4. The [[and [[<- operators allow operating on a recursive subset of a nested list.
 - But these only operate on a single recursive subset, and are not vectorized for multiple recursive subsets of a nested list at once.
 - 'squarebrackets' provides a way to reshape a nested list into a recursive matrix, thereby allowing vectorized operations on recursive subsets of such a nested list.
- 5. The [<- operator only supports copy-on-modify semantics for most classes.

 The 'squarebrackets' methods provides explicit pass-by-reference and pass-by-value semantics, whilst still respecting things like binding-locks and mutability rules.

6. 'squarebrackets' supports index-less sub-set operations, which is more memory efficient (and better for the environment) for long vectors than sub-set operations using the [and [<- operators.

Goal

Among programming languages, 'R' has perhaps one of the most flexible and comprehensive subsetting functionality, provided by the square brackets operators ([, [<-)).

But in some situations the square brackets operators are occasionally less than optimally convenient

The Goal of the 'squarebrackets' package is not to replace the square-brackets operators, but to provide **alternative** sub-setting methods and functions, to be used in situations where the square bracket operators are inconvenient.

Supported Structures

'squarebrackets' only supports the most common S3 classes, and only those that primarily use square brackets for sub-setting (hence the name of the package).

'squarebrackets' supports the following structures:

- basic atomic classes (atomic vectors, matrices, and arrays).
- mutable_atomic classes (mutable_atomic vectors, matrices, and arrays).
- factor.
- basic list classes (recursive vectors, matrices, and arrays).
- data.frame (including the classes tibble, sf-data.frame and sf-tibble).
- data.table (including the classes tidytable, sf-data.table, and sf-tidytable).

 $See\ square brackets_supported_structures\ for\ more\ details.$

Sub-set Operation Methods & Binding Implementations

The main focus of this package is on its generic methods and dimensional binding implementations.

Generic methods for atomic objects start with sb_.

Generic methods for recursive objects (list, data.frame, etc.) start with sb2_.

There is also the somewhat separate idx method, which works on both recursive and non-recursive objects.

The binding implementations for dimensional objects start with bind_.

And finally there are the slice_methods, which (currently) only work on (mutable) atomic vectors.

ACCESS SUBSETS

Methods to access subsets (i.e. extract selection, or extract all except selection):

- sb x, sb2 x: extract, exchange, or duplicate subsets.
- sb_wo, sb2_wo: return an object without the specified subset.
- sb2_rec: access recursive subsets of lists.
- slice_x: index-less and efficient, sequence-based extraction of a subset from a long vector.
- slice_wo: index-less and efficient, sequence-based returning a long vector without the specified subset.
- slicev_x: index-less and efficient, value-based extraction of a subset from a long vector.

MODIFY SUBSETS

Methods to modify subsets:

- idx: translate given indices/subscripts, for the purpose of copy-on-modify substitution.
- sb2_recin: replace, transform, remove, or add recursive subsets to a list, through R's default Copy-On-Modify semantics.
- sb_mod, sb2_mod: return the object with modified (transformed or replaced) subsets.
- Methods to rename a mutable object using pass-by-reference semantics.
- sb_set, sb2_set: modify (transform or replace) subsets of a mutable object using pass-by-reference semantics.
- slice_set: index-less and efficient, sequence-based modification of a (long) vector subset using pass-by-reference semantics.
- slicev_set: index-less and efficient, value-based modification of a (long) vector subset using pass-by-reference semantics.

EXTEND BEYOND

Methods and binding implementations, to extend or re-arrange an object beyond its current size:

- bind: implementations for binding dimensional objects.
- sb_x, sb2_x: extract, exchange, or duplicate subsets.
- sb2_recin: replace, transform, remove, or add recursive subsets to a list, through R's default Copy-On-Modify semantics.

See squarebrackets_method_dispatch for more information on how 'squarebrackets' uses its S3 Method dispatch.

Functions

SPECIALIZED FUNCTIONS

Additional specialized sub-setting functions are provided:

- lst_untree: unnest tree-like nested list into a recursive matrix, to speed-up vectorized subsetting on recursive subsets of the list.
- The dt_-functions to programmatically perform data.table-specific [-operations, with the security measures provided by the 'squarebrackets' package.
- setapply: apply functions over mutable matrix margins using pass-by-reference semantics.

HELPER FUNCTIONS

A couple of convenience functions, and helper functions for creating ranges, sequences, and indices (often needed in sub-setting) are provided:

- currentBindings: list or lock all currently existing bindings that share the share the same address as the input variable.
- n: Nested version of c, and short-hand for list.
- ndim: Get the number of dimensions of an object.
- sub2coord, coord2ind: Convert subscripts (array indices) to coordinates, coordinates to flat indices, and vice-versa.
- match_all: Find all matches, of one vector in another, taking into account the order and any duplicate values of both vectors.
- Computing indices:

idx_r to compute an integer index range.

idx_by to compute grouped indices.

idx_ord_-functions to compute ordered indices.

Overview Help Pages

Besides the website, 'squarebrackets' comes with several help pages that can be accessed from within 'R'.

MAIN DOCUMENTATION:

- squarebrackets_supported_structures:
 Lists the structures that are supported by 'squarebrackets', and explains some related terminology.
- squarebrackets_indx_fundamentals: Explains the essential fundamentals of the indexing forms in 'squarebrackets'.
- squarebrackets_indx_args:

Explains the common indexing arguments used in the main S3 methods.

• squarebrackets_modify:

Explains the essentials of modification in 'squarebrackets'

• squarebrackets_options:

Lists and explains the options the user can specify in 'squarebrackets'.

• squarebrackets_method_dispatch:

Gives details regarding the S3 method dispatch in 'squarebrackets'.

ADDITIONAL DOCUMENTATION:

• squarebrackets_PassByReference:

Explains Pass-by-Reference semantics, and its important consequences.

If you are not planning on using the pass-by-reference functionality in 'squarebrackets', you do not need to read this help page.

• squarebrackets_coercion:

Explains the difference in coercion rules between modification through Pass-by-Reference semantics and modification through copy (i.e. pass-by-value) for the supported mutable structures.

If you are not planning on using the pass-by-reference functionality in 'squarebrackets', you do not need to read this help page.

• squarebrackets_slicev:

Explains the arguments for the slicev set of methods.

If you are not planning to use the slicev methods, you do not need to read this help page.

Properties Details

The alternative sub-setting methods and functions provided by 'squarebrackets' have the following properties:

• Programmatically friendly:

- Unlike base [, it's not required to know the number of dimensions of an array a-priori, to perform subset-operations on an array.
- Missing arguments can be filled with NULL, instead of using dark magic like base::quote(expr =).
- No Non-standard evaluation.
- Functions are pipe-friendly.
- No (silent) vector recycling.
- Extracting and removing subsets uses the same syntax.

• Class consistent:

sub-setting of multi-dimensional objects by specifying dimensions (i.e. rows, columns, ...) use drop = FALSE.

So matrix in, matrix out.

The methods deliver the same results for data.frames, data.tables, tibbles, and tidytables.
 No longer does one have to re-learn the different brackets-based sub-setting rules for different types of data.frame-like objects.

Powered by the subclass agnostic 'C'-code from 'collapse' and 'data.table'.

• Explicit copy semantics:

 Sub-set operations that change its memory allocations, always return a modified (partial) copy of the object. For sub-set operations that just change values in-place (similar to the [<- methods) the user can choose a method that modifies the object by reference, or choose a method that returns a (partial) copy.

• Careful handling of names:

- Sub-setting an object by index names returns ALL matches with the given names, not just the first.
- Data.frame-like objects (see supported classes below) are forced to have unique column names.
- Sub-setting arrays using x[indx1, indx2, etc.] will drop names(x).
 The methods from 'squarebrackets' will not drop names(x).
- · Concise function and argument names.
- Performance & Energy aware:

Despite the many checks performed, the functions are kept reasonably speedy, through the use of the 'Rcpp', 'collapse', and 'data.table' R-packages.

The functions were also made to be as memory efficient as reasonably possible, to lower the carbon footprint of this package.

Author(s)

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References

The badges shown in the documentation of this R-package were made using the services of: https://shields.io/

aaa01_squarebrackets_supported_structures $Supported\ Structures$

Description

'squarebrackets' only supports the most common S3 objects, and only those that primarily use square brackets for sub-set operations (hence the name of the package).

One can generally divide the structures supported by 'squarebrackets' along 3 key properties:

• atomic vs recursive:

Types logical, integer, double, complex, character, and raw are atomic. Lists and data.frames are recursive.

• dimensionality:

Whether an object is a vector, array, or data.frame.

Note that a matrix is simply an array with 2 dimensions.

• mutability:

Base R's S3 classes (except Environments) are generally immutable:

Modifying the object will create a copy (called 'copy-on-modify').

'squarebrackets also supports data.tables and mutable_atomic objects, which are mutable:

If desired, one can modify them without copy using pass-by-reference semantics.

Supported Structures

'squarebrackets' supports the following immutable structures:

- basic atomic classes (atomic vectors and arrays).
- · factor.
- basic list classes (recursive vectors and arrays).
- data.frame
 (including the classes tibble, sf-data.frame and sf-tibble).

'squarebrackets' supports the following mutable structures:

- mutable_atomic (mutable_atomic vectors arrays);
- data.table (including the classes tidytable, sf-data.table, and sf-tidytable).

Details

Atomic vs Recursive

The sb_ methods provided by 'squarebrackets' work on **atomic** (see is.atomic) objects. The sb2_ methods provided by 'squarebrackets' work on **recursive** (see is.recursive) objects. See squarebrackets_method_dispatch for more details on the method dispatch used by 'squarebrackets'.

Dimensionality

'squarebrackets' supports dimensionless or vector objects (i.e. ndim == 0L).

squarebrackets' supports arrays (see is.array and is.matrix); note that a matrix is simply an array with 2 dimensions.

'squarebrackets' also supports data.frame-like objects (see is.data.frame).

Specifically, squarebrackets' supports a wide variety of data.frame classes:

data.frame, data.table, tibble, tidytable;

'squarebrackets' also supports their 'sf'-package compatible counter-parts:

sf-data.frame, sf-data.table, sf-tibble, sf-tidytable.

Dimensionless vectors and dimensional arrays are supported in both their atomic and recursive forms.

Data.frame-like objects, in contrast, only exist in the recursive form (and, as stated, are supported by 'squarebrackets').

Recursive vectors, recursive matrices, and recursive arrays, are collectively referred to as "lists" in the 'squarebrackets' documentation.

Note that the dimensionality of data.frame-like objects is not the same as the dimensionality of (recursive) arrays/matrices.

For example:

For any array/matrix x, it holds that length(x) == prod(dim(x)).

But for any data.frame x, it is the case that length(x) == ncol(x).

Mutable vs Immutable

Most of base R's S3 classes (except Environments) are generally immutable:

Modifying the object will create a copy (called 'copy-on-modify').

They have no explicit pass-by-reference semantics.

Most S3 objects in base 'R' are immutable:

Environments do have pass-by-reference semantics, but they are not supported by 'squarebrackets'.

Supported mutable structures:

- 'squarebrackets' supports the mutable data.table class (and thus also tidytable, which inherits from data.table).
- 'squarebrackets' also includes a new class of mutable objects: mutable_atomic objects.

mutable_atomic objects are the same as atomic objects, except they are mutable (hence the name).

Supported immutable structures:

Atomic and recursive vectors/matrices/arrays, data.frames, and tibbles.

All the functions in the 'squarebrackets' package with the word "set" in their name perform passby-reference modification, and thus only work on mutable structures.

All other functions work the same way for both mutable and immutable structures.

Derived Atomic Vector

A special class of objects are the Derived Atomic Vector structures: structures that are derived from atomic objects, but behave differently. For example:

Factors, datetime, POSIXct and so on are derived from atomic vectors.

But they have attributes and special methods that make them behave differently.

'squarebrackets' treats derived atomic classes as regular atomic vectors.

There are highly specialized packages to handle objects derived from atomic objects.

For example, the 'anytime' package to handle date-time objects.

'squarebrackets does provide some more explicit support for factors.

Not Supported S3 structures

Key-Values storage S3 structures, such as environments, are not supported by 'squarebrackets'.

Description

This help page explains the fundamentals regarding how 'squarebrackets' treats indexing. Some familiarity with base R's [and [<- operators is required to follow this help page.

Indexing Types

Base 'R' supports indexing through logical, integer, and character vectors. 'squarebrackets' supports these also (albeit with some improvements), but also supports some addi-

Whole numbers

tional methods of indexing.

Whole numbers are the most basic form on index selection.

All forms of indexing in 'squarebrackets' are internally translated to integer (or double if > (2^31 - 1)) indexing first, ensuring consistency.

Indexing through integer/numeric indices in 'squarebrackets' works the same as in base 'R', except that negative values are not allowed.

So indexing starts at 1 and is inclusive.

Logical

Selecting indices with a logical vector in 'squarebrackets' works the same as in base 'R', except that recycling is not allowed.

Characters

When selecting indices using a character vector, base 'R' only selects the first matches in the names. 'squarebrackets', however, selects all matches:

```
nms <- c("a", letters[4:1], letters[1:5])
x <- 1:10
names(x) <- nms
print(x) #' `x` has multiple elements with the name "a"
#> a d c b a a b c d e
#> 1 2 3 4 5 6 7 8 9 10

sb_x(x, "a") # extracts all indices with the name "a"
#> a a
#> 1 5 6

sb_x(x, c("a", "a")) # repeats all indices with the name "a"
```

```
#> a a a a a a
#> 1 5 6 1 5 6
```

Character indices are internally translated to integer indices using match_all.

Imaginary Numbers

A complex vector y is structured as

```
y = a + b * i
```

where Re(y) returns a, and Im(y) returns b.

squarebrackets' includes support for indexing through imaginary numbers (Im(y)) of complex vectors.

Indexing with imaginary numbers is a generalization of indexing with regular integers.

It works as follows:

Imaginary numbers that are positive integers, like $1:10 \times 1i$, work the same as regular integers. Imaginary numbers that are negative integers, like $1:10 \times -1i$, index by counting backwards (i.e. from the end).

Note that **only** the Imaginary part of a complex vector is used (Im(y)); the Real part (Re(y)) is **ignored**.

See the results of the following code as an example:

```
x <- 1:30 # vector of 30 elements

sb_x(x, 1:10 * 1i) # extract first 10 elements
#> [1] 1 2 3 4 5 6 7 8 9 10

sb_x(x, 1:10 * -1i) # extract last 10 elements
#> [1] 30 29 28 27 26 25 24 23 22 21

sb_x(x, 10:1 * -1i) # last 10 elements, in tail()-like order
#> [1] 21 22 23 24 25 26 27 28 29 30
```

Thus complex vectors allow the user to choose between counting from the beginning, like regular integers, or backwards counting from the end.

Flat Indices and Subscripts

The primary indexing argument for vectors (i.e. dimensionless objects), is the i argument, which represents flat indices.

The primary indexing argument for dimensional objects supported by 'squarebrackets' (i.e. arrays and data.frame-like objects), is the s, d argument pair, which represent "subscripts".

(Given, for example, a 3-dimensional array, the subscript [1:10, 2:5, 3:9], refers to rows 1 to 10, columns 2 to 5, and layers 3 to 9.)

This s, d argument pair works consistently for any dimensional object supported by 'squarebrackets', and does not require a-priori knowledge on the number of dimensions the object has.

This is particularly useful for arrays, which can have any number of dimensions.

Arrays and matrices (matrices are simply arrays with 2 dimensions) support both flat indices and subscripts.

In that case the flat indices, also called linear indices, specify the indices of an array as-if it is vector, thus ignoring dimensions.

For the relationship between flat indices and subscripts for arrays, see the sub2ind help page.

Inverting

Inverting indices means to specify all elements **except** the given indices.

Consider for example the atomic vector month. abb (abbreviate month names).

Given this vector, indices 1:5 gives c("Jan" "Feb" "Mar" "Apr", "May").

Inverting those same indices will give c("Jun" "Jul" "Aug" "Sep" "Oct" "Nov" "Dec").

In base 'R', inverting an index is done in different ways.

(negative numbers for numeric indexing, negation for logical indexing, manually un-matching for character vectors).

'squarebrackets' provides a (somewhat) consistent syntax to invert indices:

- The methods that end with _x perform extraction; to invert extraction, i.e. return the object **without** the specified subset, use the methods that end with _wo.
- In the modification methods (_mod_/_set_) one can set the argument inv = TRUE to invert indices.

EXAMPLES

```
x <- month.abb
print(x)
#> [1] "Jan" "Feb" "Mar" "Apr" "May" "Jun" "Jul" "Aug" "Sep" "Oct" "Nov" "Dec"

sb_x(x, 1:5) # extract first 5 elements
#> [1] "Jan" "Feb" "Mar" "Apr" "May"

sb_wo(x, 1:5) # return WITHOUT first 5 elements
#> [1] "Jun" "Jul" "Aug" "Sep" "Oct" "Nov" "Dec"

sb_mod(x, 1:5, rp = "XXX") # copy, replace first 5 elements, return result
#> [1] "XXX" "XXX" "XXX" "XXX" "XXX" "Jun" "Jul" "Aug" "Sep" "Oct" "Nov" "Dec"

sb_mod(x, 1:5, inv = T, rp = "XXX") # same, but for all except first 5 elements
#> [1] "Jan" "Feb" "Mar" "Apr" "May" "XXX" "XXX" "XXX" "XXX" "XXX" "XXX" "XXX" "XXXX" "XXX"
```

ABOUT ORDERING

The order in which the user gives indices when inverting indices generally does not matter.

The order of the indices as they appear in the original object x is maintained, just like in base 'R'.

Out-of-Bounds Integers, Non-Existing Names, and NAs

- Integer indices that are out of bounds (including NaN and NA_integer_) always give an error.
- Character indices that specify non-existing names is considered a form of zero-length indexing.

Specifying NA names returns an error.

• Logical indices are translated internally to integers using which, and so NAs are ignored.

Index-less Sub-set Operations

Until now this help page focussed on performing sub-set operations with an indexing vector.

Performing sub-set operations on a long vector using a index vector (which may itself also be a long vector) is not very memory-efficient.

'squarebrackets' therefore introduces index-less sub-set operations, through the slice_ and slicev_ methods.

These methods are much more memory and computationally efficient than index-based sub-set methods (and so also a bit better for the environment!).

The slice_ methods perform sequence based sub-set operations.

The slicev_ methods (notice the "v" at the end) perform value-based sub-set operations.

Though this method is intentionally kept relatively simple, it is still involved enough to warrant its own help page;

for the details on value-based index-less sub-set operations, please see squarebrackets_slicev.

Regarding Performance

Integer vectors created through the : operator are "compact ALTREP" integer vectors, and provide the fastest way to specify indices.

Indexing through names (i.e. character vectors) is the slowest.

Complex vectors of imaginary numbers are somewhat in the middle in terms of speed.

Index-less sub-set operations are usually faster and more memory efficient than any index-based sub-set operation.

So if performance is important, use index-less sub-set operations, or use compact ALTREP integer indices.

Indexing in Recursive Subsets

Until now this help page focussed on indexing for regular (or "shallow") subsets.

This section will discuss indexing in recursive subsets.

One of the differences between atomic and recursive objects, is that recursive objects support recursive subsets, while atomic objects do not.

Bear in mind that every element in a recursive object is a reference to another object. Consider the following list x:

```
x <- list(
    A = 1:10,
    B = letters,
    C = list(A = 11:20, B = month.abb)
)</pre>
```

Regular subsets, AKA surface-level subset operations ([, [<- in base 'R'), operate on the recursive object itself.

```
I.e. sb2_x(x, 1), or equivalently x[1], returns the list list(A = 1:10):
```

```
sb2_x(x, 1) # equivalent to x[1]; returns list(A = 1:10) #> $A #> [1] 1 2 3 4 5 6 7 8 9 10
```

Recursive subset operations ([[, [[<-, and \$ in base 'R'), on the other hand, operate on an object a subset of the recursive object references to.

I.e. $sb2_rec(x, 1)$, or equivalently x[[1]], returns the **integer vector** 1:10:

```
sb2_rec(x, 1) # equivalent to x[[1]]; returns 1:10 #> [1] 1 2 3 4 5 6 7 8 9 10
```

Recursive objects can refer to other recursive objects, which can themselves refer to recursive objects, and so on.

Recursive subsets can go however deep you want.

So, for example, to extract the character vector month.abb from the aforementioned list x, one would need to do:

```
sb2_rec(x, c("C","B")), (in base R: x$C$B):
sb2_rec(x, c("C","B")) # equivalent to x$C$B
#> [1] "Jan" "Feb" "Mar" "Apr" "May" "Jun" "Jul" "Aug" "Sep" "Oct" "Nov" "Dec"
# or:
sb2_rec(x, c(3, 2)) # equivalent to x[[3]][[2]]
#> [1] "Jan" "Feb" "Mar" "Apr" "May" "Jun" "Jul" "Aug" "Sep" "Oct" "Nov" "Dec"
```

LIMITATIONS

Indexing in recursive subsets is significantly more limited than in regular (or "shallow") subsets:

• Recursive subset operations using sb2_rec/sb2_recin only support positive integer vectors and character vectors.

- Imaginary numbers (using complex vectors) and logical vectors are not supported.
- Since a recursive subset operation only operates on a single element, specifying the index with a character vector only selects the first matching element (just like base 'R'), not all matches.
- Inverting indices is also **not** available for recursive indexing.
- Unlike regular sub-setting, out-of-bounds specification for indices is acceptable, as it can be used to add new values to lists.

Non-Standard Evaluation

'squarebrackets' is designed primarily for programming, and seeks to be fully programmatically friendly.

As part of this endeavour, 'squarebrackets' never uses Non-Standard Evaluation.

All input for all methods and functions in 'squarebrackets' are objects that can be stored in a variable.

Like atomic vectors, lists, formulas, etc.

aaa03_squarebrackets_indx_args

Index Arguments in the Generic Sub-setting Methods

Description

There are several types of arguments that can be used in the generic methods of 'squarebrackets' to specify the indices to perform operations on:

- i: to specify flat (i.e. dimensionless) indices.
- s, d: to specify indices of arbitrary dimensions in any dimensional object supported by 'squarebrackets' (i.e. arrays and data.frame-like objects).
- margin, slice: to specify indices of one particular dimension (for arrays and data.frame-like objects).

Only used in the idx method.

• obs, vars: to specify observations and/or variables in specifically in data.frame-like objects.

For the fundamentals of indexing in 'squarebrackets', see squarebrackets_indx_fundamentals. In this help page x refers to the object on which subset operations are performed.

Argument i

class: atomic vector class: derived atomic vector class: recursive vector class: atomic array class: recursive array

Any of the following can be specified for argument i:

- NULL, corresponds to missing argument.
- a vector of length 0, in which case no indices are selected for the operation (i.e. empty selection).
- a numeric vector of **strictly positive whole numbers** giving indices.
- a complex vector, as explained in squarebrackets indx fundamentals.
- a **logical vector**, of the same length as x, giving the indices to select for the operation.
- a **character** vector of index names.

 If an object has multiple indices with the given name, ALL the corrections of the correction of th

If an object has multiple indices with the given name, ALL the corresponding indices will be selected for the operation.

• a **function** that takes as input x, and returns a logical vector, giving the element indices to select for the operation.

For atomic objects, i is interpreted as i(x).

For recursive objects, i is interpreted as lapply(x, i).

Using the i arguments corresponds to doing something like the following:

```
sb_x(x, i = i) # ==> x[i] # if `x` is atomic 
 <math>sb_x(x, i = i) # ==> x[i] # if `x` is recursive
```

If i is a function, it corresponds to the following:

```
sb_x(x, i = i) \# ==> x[i(x)] \# if `x` is atomic 
 <math>sb_x(x, i = i) \# ==> x[lapply(x, i)] \# if `x` is recursive
```

Argument Pair s, d

class: atomic array class: recursive array class: data.frame-like

The s, d argument pair, inspired by the abind::asub function from the 'abind' package, is the primary indexing argument for sub-set operations on dimensional objects.

The s argument specifies the **subscripts** (i.e. dimensional indices).

The d argument gives the dimensions for which the s holds (i.e. d specifies the "non-missing" margins).

The d argument must be an integer vector.

s must be a list of length 1, or a list of the same length as d. If s is a list of length 1, it is internally recycled to become the same length as d.

Each element of s can be any of the following:

- a vector of length 0, in which case no indices are selected for the operation (i.e. empty selection).
- a numeric vector of **strictly positive whole numbers** with indices of the specified dimension to select for the operation.
- a **complex** vector, as explained in squarebrackets_indx_fundamentals.
- a **logical** vector of the same length as the corresponding dimension size, giving the indices of the specified dimension to select for the operation.
- a **character** vector giving the dimnames to select.

 If a dimension has multiple indices with the given name, ALL the corresponding indices will be selected for the operation.

Note the following:

- As stated, d specifies which index margins are non-missing. If d is of length 0, it is taken as "all index margins are missing".
- The default value for d is 1:ndim(x).

To keep the syntax short, the user can use the n function instead of list() to specify s.

EXAMPLES

Here are some examples for clarity, using an atomic array x of 3 dimensions:

- sb_x(x, n(1:10, 1:5), c(1, 3)) extracts the first 10 rows, all columns, and the first 5 layers, of array x.
- sb_x(x, n(1:10), 2) extracts the first 10 columns of array x.
- sb_x(x, n(1:10)), extracts the first 10 rows, columns, and layers of array x.
- sb_x(x, n(1:10), c(1, 3)), extracts the first 10 rows, all columns, and the first 10 layers, of array x.

I.e.:

```
sb_x(x, n(1:10, 1:5), c(1, 3)) # ==> x[1:10, , 1:5, drop = FALSE]
sb_x(x, n(1:10), 2) # ==> x[ , 1:10, , drop = FALSE]
sb_x(x, n(1:10)) # ==> x[1:10, 1:10, drop = FALSE]
sb_x(x, n(1:10), c(1, 3)) # ==> x[1:10, , 1:10, drop = FALSE]
```

NOTE

If length(d) is 1, s can also be given as an atomic vector (of any length), instead of a list of length

Although it is allowed for s and d to both be atomic vectors of length 1, for the readability of your code it is **highly recommended** that s and d be explicitly **named** in your method call, in such a case.

I.e.:

```
sb_x(x, 1, 1) \# BAD: this is not very readable sb_x(x, s = 1, d = 1) \# This is GOOD
```

For a brief explanation of the relationship between flat indices (i) and subscripts (s, d) in arrays, see sub2ind.

Argument Pair margin, slice

class: atomic array class: recursive array class: data.frame-like

Relevant only for the idx method.

The margin argument specifies the dimension on which argument slice is used.

I.e. when margin = 1, slice selects rows; when margin = 2, slice selects columns; etc.

The slice argument can be any of the following:

- a numeric vector of **strictly positive whole numbers** with dimension indices to select for the operation.
- a **complex** vector, as explained in squarebrackets_indx_fundamentals.
- a **logical** vector of the same length as the corresponding dimension size, giving the dimension indices to select for the operation.
- a **character** vector of index names.

If a dimension has multiple indices with the given name, ALL the corresponding indices will be selected for the operation.

One could also give a vector of length \emptyset for slice;

Argument slice is only used in the idx method , and the result of idx are meant to be used inside the regular [and [<- operators.

Thus the effect of a zero-length index specification depends on the rule-set of [.class(x)] and [<-.class(x)].

Arguments obs, vars

class: data.frame-like

The obs argument specifies indices for observations (i.e. rows) in data.frame-like objects.

The vars argument specifies indices for variables (i.e. columns) in data.frame-like objects.

The obs and vars arguments are inspired by the subset and select arguments, respectively, of base R's subset.data.frame method. However, the obs and vars arguments do **not** use non-standard evaluation, as to keep 'squarebrackets' fully programmatically friendly.

The obs Argument

The obs argument can be any of the following:

- NULL (default), corresponds to a missing argument.
- a vector of length 0, in which case no indices are selected for the operation (i.e. empty selection).
- a numeric vector of strictly positive whole numbers with row indices to select for the operation.
- a **complex** vector, as explained in squarebrackets_indx_fundamentals.
- a **logical** vector of the same length as the number of rows, giving the row indices to select for the operation.
- a **one-sided formula**, with a single logical expression using the column names of the data.frame, giving the condition which observation/row indices should be selected for the operation.

So to perform an operation on the observations for which holds that height > 2 and sex != "female", specify the following formula:

```
obs = ~ (height > 2) & (sex != "female")
```

If the formula is linked to an environment, any variables not found in the data set will be searched from the environment.

The vars Argument

The vars argument can be any of the following

- NULL (default), corresponds to a missing argument.
- a vector of length 0, in which case no indices are selected for the operation (i.e. empty selection).
- a numeric vector of **strictly positive whole numbers** with column indices to select for the operation.
- a **complex** vector, as explained in squarebrackets_indx_fundamentals.
- a **logical** vector of the same length as the number of columns, giving the column indices to select for the operation.
- a **character** vector giving the colnamess to select.

 Note that 'squarebrackets' assumes data.frame-like objects have unique column names.
- a **function** that returns a logical vector, giving the column indices to select for the operation. For example, to select all numeric variables, specify vars = is.numeric.
- a **two-sided formula**, where each side consists of a single term, giving a range of names to select

For example, to select all variables between and including the variables "height" and "weight", specify the following:

```
vars = heigth ~ weight.
```

EXAMPLE

So using the obs, vars arguments corresponds to doing something like the following:

```
sb2_x(x, obs = obs, vars = vars) # ==> subset(x, ...obs..., ...vars...)
```

Argument inv

all classes

Relevant for the sb mod/sb2 mod, sb set/sb2 set, and idx methods.

By default, inv = FALSE, which translates the indices like normally.

When inv = TRUE, the inverse of the indices is taken.

Consider, for example, an atomic matrix x;

using $sb_mod(x, 1:2, 2L, tf = tf)$ corresponds to something like the following:

```
x[, 1:2] <- tf(x[, 1:2])
x
```

and using $sb_mod(x, vars = 1:2, inv = TRUE, tf = tf)$ corresponds to something like the following:

```
x[, -1:-2] \leftarrow tf(x[, -1:-2])
```

NOTE

The order in which the user gives indices when inv = TRUE generally does not matter.

The order of the indices as they appear in the original object x is maintained, just like in base 'R'. Therefore, when replacing multiple values where the order of the replacement matters, it is better to keep inv = FALSE, which is the default.

For replacement with a single value or with a transformation function, inv = TRUE can be used without considering the ordering.

All Missing Indices

NULL in the indexing arguments corresponds to a missing argument.

For s, d, specifying d of length 0 also corresponds to all subscripts being missing.

Thus, for **both** sb_x/sb2_x and sb_wo/sb2_wo, using missing or NULL indexing arguments for all indexing arguments corresponds to something like the following:

x[]

Similarly, for sb_mod/sb2_mod and sb_set/sb2_set, using missing or NULL indexing arguments corresponds to something like the following:

```
x[] <- rp # for replacement
x[] <- tf(x) # for transformation</pre>
```

The above is true **even if** inv = TRUE and/or red = TRUE.

Disallowed Combinations of Index Arguments

One cannot specify i and the other indexing arguments simultaneously; it's either i, or the other arguments.

One cannot specify row and filter simultaneously; it's either one or the other.

One cannot specify col and vars simultaneously; it's either one or the other.

One cannot specify the s, d pair and slice, margin pair simultaneously; it's either one pair or the other pair.

In the above cases it holds that if one set is specified, the other is set is ignored.

Drop

Sub-setting with the generic methods from the 'squarebrackets' R-package using dimensional arguments (s, d, row, col filter, vars) always use drop = FALSE.

To drop potentially redundant (i.e. single level) dimensions, use the drop function, like so:

$$sb_x(x, s, d) > drop() # ==> x[..., drop = TRUE]$$

References

Plate T, Heiberger R (2016). *abind: Combine Multidimensional Arrays*. R package version 1.4-5, https://CRAN.R-project.org/package=abind.

```
aaa04\_squarebrackets\_modify Regarding Modification
```

Description

This help page describes the main modification semantics available in 'squarebrackets'.

Base R's default modification

For most average users, R's default copy-on-modify semantics are fine.

The benefits of the indexing arguments from 'squarebrackets' can be combined the [<- operator, through the idx method.

The result of the idx() method can be used inside the regular square-brackets operators.

For example like so:

```
x <- array(...)
my_indices <- idx(x, s, d)
x[my_indices] <- value

y <- data.frame(...)
rows <- idx(y, 1:10, 1, inv = TRUE)
cols <- idx(y, c("a", "b"), 2)
y[rows, cols] <- value</pre>
```

thus allowing the user to benefit from the convenient index translations from 'squarebrackets', whilst still using R's default copy-on-modification semantics (instead of the semantics provided by 'squarebrackets').

Explicit Copy

'squarebrackets' provides the sb_mod/sb2_mod method to modify through a (shallow) copy. This method returns the modified object.

For recursive objects, sb2_mod returns the original object, where only the modified subsets are copied, thus preventing unnecessary usage of memory.

Pass-by-Reference

'squarebrackets' provides the sb_set/sb2_set and slice_set methods to modify by reference, meaning no copy is made at all.

Pass-by-Reference is fastest and the most memory efficient.

But it is also more involved than the other modification forms, and requires more thought.

See squarebrackets_PassByReference for more information.

Replacement and Transformation in Atomic Objects

The rp argument is used to replace the values at the specified indices with the values specified in rp. Using the rp argument in the modification methods, corresponds to something like the following:

```
x[...] <- rp
```

The tf argument is used to transform the values at the specified indices through transformation function tf. Using the tf argument corresponds to something like the following:

```
x[...] \leftarrow tf(x[...])
```

where tf is a function that **returns** an object of appropriate type and size (so tf should not be a pass-by-reference function).

Replacement and Transformation in Lists

The rp and tf arguments work mostly in the same way for recursive objects. But there are some slight differences.

Argument rp

'squarebrackets' demands that rp is always provided as a list in the S3 methods for recursive vectors, matrices, and arrays (i.e. lists).

This is to prevent ambiguity with respect to how the replacement is recycled or distributed over the specified indices

(See Footnote 1 below).

Argument tf

Most functions in (base) 'R' are vectorized for atomic objects, but not for lists (see Footnote 2 below).

'squarebrackets' will therefore apply transformation function tf via lapply, like so:

$$x[...] \leftarrow lapply(x[...], tf)$$

In the methods for recursive objects, the tf argument is accompanied by the .lapply argument. By default, .lapply = lapply.

The user may supply a custom lapply()-like function in this argument to use instead.

For example, to perform parallel transformation, the user may supply future.apply::future_lapply. The supplied function must use the exact same argument convention as lapply, otherwise errors or unexpected behaviour may occur.

Replacement and Transformation in data.frame-like Objects

Replacement and transformations in data.frame-like objects are a bit more flexible than in Lists.

rp is not always demanded to be a list for data.frame-like objects, only when appropriate (for example, when replacing multiple columns, or when the column itself is a list.)

When rp is given as a list, it is unclassed and unnamed before being used to replace values.

This is to ensure consistency across all supported data.frame types.

Bear in mind that every column in a data.frame is like an element in a list; so .lapply is used for transformations across multiple columns.

Recycling and Coercion

Recycling is not allowed in the modification methods.

So, for example, length(rp) must be equal to the length of the selected subset, or equal to 1.

When using Pass-by-Reference semantics, the user should be extra mindful of the auto-coercion rules

See squarebrackets_coercion for details.

Footnotes

Footnote 1

Consider the following replacement in base 'R':

```
x <-list(1, 2, 3, 4, 5, 6, 7, 8, 9, 10)
x[1:2] <- 2:1
```

What will happen?

Will the x[1] be list(1:2) and x[2] also be list(1:2)?

Or will x[1] be list(2) and x[2] be list(1)?

It turns out the latter will happen; but this is somewhat ambiguous from the code.

To prevent such ambiguity in your code, 'squarebrackets' demands that rp is always provided as a list.

Footnote 2

Most functions in (base) 'R' are vectorized for atomic objects, but not for lists.

One of the reasons is the following:

In an atomic vector x of some type t, every single element of x is a scalar of type t.

However, every element of some list x can be virtually anything:

an atomic object, another list, an unevaluated expression, even dark magic like quote(expr =).

It is difficult to make a vectorized function for an object with so many unknowns.

Therefore, in the vast majority of the cases, one needs to loop through the list elements.

 ${\tt aaa05_squarebrackets_options}$

squarebrackets Options

Description

This help page explains the various global options that can be set for the 'squarebrackets' package, and how it affects the functionality.

Check Duplicates

argument: chkdup

option: squarebrackets.chkdup

The sb_x/sb2_x methods are the only methods where providing duplicate indices actually make sense.

For the other methods, it doesn't make sense.

Giving duplicate indices usually won't break anything; however, when replacing/transforming or removing subsets, it is almost certainly not the intention to provide duplicate indices.

Providing duplicate indices anyway might lead to unexpected results.

Therefore, for the methods where giving duplicate indices does not make sense, the chkdup argument is present.

This argument controls whether the method in question checks for duplicates (TRUE) or not (FALSE).

Setting chkdup = TRUE means the method in question will check for duplicate indices, and give an error when it finds them.

Setting chkdup = FALSE will disable these checks, which saves time and computation power, and is thus more efficient.

Since checking for duplicates can be expensive, it is set to FALSE by default.

The default can be changed in the squarebrackets.chkdup option.

Sticky

argument: sticky

option: squarebrackets.sticky

The slice_x, slice_wo, and slicev_x methods can already handle names (through the use.names argument), attributes specific to the mutable_atomic class, and attributes specific to the factor class.

Attributes which are not names, and not specific to mutable_atomic class, and not specific to the factor class - henceforth referred to as "other attributes" - are treated differently.

How the slice_x, slice_wo, and slicev_x methods handle these "other" attributes, is determined by the sticky option and argument.

When sticky = FALSE, the slice_x, slice_wo, and slicev_x methods will drop all **other** attributes.

By setting sticky = TRUE, all these **other** attributes, except comment and tsp, will be preserved; The key advantage for this, is that classes that use static attributes (i.e. classes that use attributes that do not change when sub-setting), are automatically supported if sticky = TRUE, and no separate methods have to written for the slice_x, slice_wo, and slicev_x methods.

Attributes specific to classes like difftime, Date, POSIXct, roman, hexmode, octmode, and more, use static attributes.

Instead of setting sticky = TRUE or sticky = FALSE, one can also specify all classes that use static attributes that you'll be using in the current R session.

In fact, when 'squarebrackets' is **loaded**, the squarebrackets.sticky option is set as follows:

```
squarebrackets.sticky = c(
   "difftime", "Date", "POSIXct", "roman", "hexmode", "octmode"
```

)

```
So in the above default setting, sticky = TRUE for "difftime", "Date", "POSIXct", "roman", "hexmode", "octmode".

Also in the above default setting, sticky = FALSE for other classes.
```

The reason the slice_x, slice_wo, and slicev_x methods need the sticky option, is because of the following.

Unlike most sb_/sb2_ methods, the slice_x, slice_wo, and slicev_x methods are not wrappers around the [and [<- operators.

Therefore, most [- S3 methods for highly specialized classes are not readily available for the slice x, slice wo, and slicev x methods.

Which in turn means important class-specific attributes are not automatically preserved.

The sticky option is a convenient way to support a large number of classes, without having to write specific methods for them.

For specialized classes that use attributes that **do** change when sub-setting, separate dispatches for the slice_x, slice_wo, and slicev_x methods need to be written.

Package authors are welcome to create method dispatches for their own classes for these methods.

Description

This help page gives some additional details regarding the S3 method dispatch used in 'squarebrackets'.

Atomic vs Recursive

Atomic and recursive objects are quite different from each other in some ways:

- homo- or heterogeneous: an atomic object can only have values of one data type. recursive objects can hold values of any combination of data types.
- nesting: Recursive objects can be nested, while atomic objects cannot be nested.
- **copy and coercion effect**: One can coerce or copy a subset of a recursive object, without copying the rest of the object.
 - For atomic objects, however, a coercion or copy operation coerces or copies the entire vector (ignoring attributes).
- **vectorization**: most vectorized operations generally work on atomic objects, whereas recursive objects often require loops or apply-like functions.

• **recursive subsets**: Recursive objects distinguish between "regular" subset operations (in base R using [, [<-), and recursive subset operations (in base R using [[, [[<-).

See for example the sb2_rec method, or the red = TRUE argument in the sb2_x and sb2_wo methods.

For atomic objects, these 2 have no meaningful difference (safe for perhaps some minor attribute handling).

• **views**: For recursive objects, one can create a view of a recursive subset. Subset views do not exist for atomic objects.

Despite these non-trivial differences, the S3 method dispatch does not distinguish between atomic and recursive objects.

I.e. S3 methods check if an object is, for example, an array, but not if it is an atomic array or a recursive array.

(S3 method dispatch actually does distinguish between basic atomic and recursive vectors, but not for dimensional objects like arrays, which is problematic for this specific package).

Therefore, the methods in 'squarebrackets' that perform subset operations on an object, come in the atomic (sb_) and recursive (sb2_) form.

The idx method operates on the indices of an object, but does not operate on the object itself, and so has no distinction between the atomic and recursive form.

Manual Dispatch

The 'squarebrackets' package intentionally exports each function in its S3 method dispatch system. This is handy for programming purposes.

For example: one can explicitly alias a specific dispatch of a method, if one so desires.

For example like so:

```
array_x <- function(x, ...) {
   if(is.atomic(x)) {
     sb_x.array(x, ...)
   }
   else if(is.recursive(x)) {
     sb2_x.array(x, ...)
   }
}</pre>
```

Under certain circumstances, this might help your code to be more clear.

Ellipsis

Due to how the S3 method dispatch system works in 'R', all generic methods have the ellipsis argument (...).

For the user's safety, 'squarebrackets' does check that the user doesn't accidentally add arguments that make no sense for that method (like specifying the inv argument when calling sb_x).

```
aaa07_squarebrackets_PassByReference

Regarding Modification By Reference
```

Description

This help page describes how modification using "pass-by-reference" semantics is handled by the 'squarebrackets' package.

This help page does not explain all the basics of pass-by-reference semantics, as this is treated as prior knowledge.

All functions/methods in the 'squarebrackets' package with the word "set" in the name use pass-by-reference semantics.

Advantages and Disadvantages

The main advantage of pass-by-reference is that much less memory is required to modify objects, and modification is also generally faster.

But it does have several disadvantages.

First, the coercion rules are slightly different: see squarebrackets_coercion.

Second, if 2 or more variables refer to exactly the same object, changing one variable also changes the other ones.

I.e. the following code,

```
x <- y <- mutable_atomic(1:16)
sb_set(x, i = 1:6, rp = 8)</pre>
```

modifies not just x, but also y.

This is true even if one of the variables is locked (see bindingIsLocked).

I.e. the following code,

```
x <- mutable_atomic(1:16)
y <- x
lockBinding("y", environment())
sb_set(x, i = 1:6, rp = 8)</pre>
```

modifies both x and y without error, even though y is a locked constant.

Mutable vs Immutable Classes

With the exception of environments, most of base R's S3 classes are treated as immutable: Modifying an object in 'R' will make a copy of the object, something called 'copy-on-modify' semantics.

A prominent mutable S3 class is the data.table class, which is a mutable data.frame class, and supported by 'squarebrackets'.

Similarly, 'squarebrackets' adds a class for mutable atomic objects: mutable_atomic.

Material vs Immaterial objects

Most objects in 'R' are material objects:

the values an object contains are actually stored in memory.

For example, given x < -rnorm(1e6), x is a material object:

1 million values (decimal numbers, in this case) are actually stored in memory.

In contrast, ActiveBindings are immaterial:

They are objects that, when accessed, call a function to generate values on the fly, rather than actually storing values.

Since immaterial objects do not actually store the values in memory, the values obviously also cannot be changed in memory.

Therefore, Pass-by-Reference semantics don't work on immaterial objects.

ALTREP

The mutable_atomic constructors (i.e. mutable_atomic, as.mutable_atomic, etc.) will automatically materialize ALTREP objects, to ensure consistent behaviour for 'pass-by-reference' semantics.

A data.table can have ALTREP columns.

A data. tables will coerce the column to a materialized column when it is modified, even by reference.

Mutability Rules With Respect To Recursive Objects

Lists are difficult objects in that they do not contain elements, they simply point to other objects, that one can access via a list.

When a recursive object is of a mutable class, all its subsets are treated as mutable, as long as they are part of the object.

On the other hand, When a recursive object is of an immutable class, its recursive subsets retain their original mutability.

Example 1: Mutable data.tables

A data. table is a mutable class.

So all columns of the data. table are treated as mutable;

There is no requirement to, for instance, first change all columns into the class of mutable_atomic

to modify these columns by reference.

Example 2: Immutable lists

A regular list is an immutable class.

So the list itself is immutable, but the recursive subsets of the list retain their mutability.

If you have a list of data. table objects, for example, the data.tables themselves remain mutable.

Therefore, the following pass-by-reference modification will work without issue:

```
x <- list(
  a = data.table::data.table(cola = 1:10, colb = letters[1:10]),
  b = data.table::data.table(cola = 11:20, colb = letters[11:20]))
myref <- x$a
sb2_set(myref, vars = "cola", tf = \(x)x^2)</pre>
```

Notice in the above code that myref has the same address as x\$a, and is therefore not a copy of x\$a. Thus changing myref also changes x\$a.

In other words: myref is what could be called a "View" of x\$a.

Input Variable

Methods/functions that perform in-place modification by reference only works on objects that actually exist as an actual variable, similar to functions in the style of $some_function(x, ...) < value$.

```
Thus things like any of the following, sb_set(1:10, ...), sb2_set(x$a, ...), or sb_set(base::letters), will not work.
```

Lock Binding

Mutable classes are, as the name suggests, meant to be mutable.

Locking the binding of a mutable object is **mostly** fruitless (but not completely; see the current-Bindings function).

To ensure an object cannot be modified by any of the methods/functions from 'squarebrackets', 2 things must be true:

- the object must be an immutable class.
- the binding must be **locked** (see lockBinding).

Protection

Due to the properties described above in this help page, 'squarebrackets' protects the user from do something like the following:

```
# letters = base::letters
sb_set(letters, i = 1, rp = "XXX")
```

'squarebrackets' will give an error when running the code above, because:

- 1. most addresses in baseenv() are protected;
- 2. immutable objects are disallowed (you'll have to create a mutable object, which will create a copy of the original, thus keeping the original object safe from modification by reference);
- 3. locked bindings are disallowed.

Examples

```
# the following code demonstrates how locked bindings,
# such as `base::letters`,
# are being safe-guarded
x <- list(a = base::letters)</pre>
myref <- x$a # view of a list
address(myref) == address(base::letters) # TRUE: point to the same memory
bindingIsLocked("letters", baseenv()) # base::letters is locked ...
bindingIsLocked("myref", environment()) # ... but this pointer is not!
if(requireNamespace("tinytest")) {
  tinytest::expect_error(
    sb_set(myref, i = 1, rp = "XXX") # this still gives an error though ...
  )
}
is.mutable_atomic(myref) # ... because it's not of class `mutable_atomic`
x <- list(
 a = as.mutable_atomic(base::letters) # `as.mutable_atomic()` makes a copy
myref <- x$a # view of a list</pre>
address(myref) == address(base::letters) # FALSE: it's a copy
 myref, i = 1, rp = "XXX" # modifies x, does NOT modify `base::letters`
print(x) # x is modified
base::letters # but this still the same
```

Description

This help page describes the auto-coercion rules of the mutable classes, as they are handled by the 'squarebrackets' package.

This useful information for users who wish to intend to employ Pass-by-Reference semantics as provided by 'squarebrackets'.

mutable atomic

```
coercion_through_copy: YES coercion_by_reference: NO
```

Mutable atomic objects are automatically coerced to fit the modified subset values, when modifying through copy, just like regular atomic classes.

For example, replacing one or multiple values in an integer vector (type int) with a decimal number (type dbl) will coerce the entire vector to type dbl.

Replacing or transforming subsets of mutable atomic objects **by reference** does not support coercion. Thus, for example, the following code,

```
x <- mutable_atomic(1:16)
sb_set(x, i = 1:6, rp = 8.5)
#> coercing replacement to integer
print(x)
#> [1] 8 8 8 8 8 8 8 7 8 9 10 11 12 13 14 15 16
#> mutable_atomic
#> typeof: integer
```

gives c(rep(8, 6) 7:16) instead of c(rep(8.5, 6), 7:16), because x is of type integer, so rp is interpreted as type integer also.

data.table, when replacing/transforming whole columns

```
coercion_through_copy: YES
coercion_by_reference: YES
```

A data table is actually a list made mutable, where each column is itself a list. As such, replacing/transforming whole columns, without specifying rows (not even 1:nrow(x)), allows completely changing the type of the column.

data.table, when partially replacing/transforming columns

```
coercion_through_copy: YES coercion_by_reference: NO
```

If rows are specified in the sb2_set method, and thus not whole columns but parts of columns are replaced or transformed, no auto-coercion takes place.

I.e.: replacing/transforming a value in an integer (int) column to become 1.5, will not coerce the column to the decimal type (db1); instead, the replacement value 1.5 is coerced to integer 1.

The sb2_mod method, however, allows for coercion just like regular data.frame objects.

Views of Lists

Regular lists are treated as immutable by 'squarebrackets'.

But remember that a list is a (potentially hierarchical) structure of references to other objects.

Thus, even if a list itself is not treated as mutable, subsets of a list which are themselves mutable classes, are mutable.

For example, if you have a list of data.table objects, the data.tables themselves are mutable.

Therefore, the following will work:

```
x <- list(
  a = data.table::data.table(cola = 1:10, colb = letters[1:10]),
  b = data.table::data.table(cola = 11:20, colb = letters[11:20]))
myref <- x$a
sb2_set(myref, vars = "cola", tf = \(x)x^2)</pre>
```

Notice in the above code that myref is not a copy of x\$a, since they have the same address.

Thus changing myref also changes x\$a.

In other words: myref is what could be called a "view" of x\$a.

Notice also that sb2_set(x\$a, ...) will not work, since sb_set/sb2_set requires actual variables, similar to in-place functions in the style of `myfun()<-`.

The auto-coercion rules of Views of Lists, depends entirely on the object itself.

Thus if the View is a data.table, coercion rules of data.tables apply.

And if the View is a mutable_atomic matrix, coercion rules of mutable_atomic matrices apply, etc.

Examples

```
str(obj) # notice that columns "a" and "c" are INTEGER (`int`)
sb2_mod(
 obj, vars = is.numeric,
 tf = sqrt # SAFE: row=NULL & obs = NULL, so coercion performed
# sb_set():
sb2_set(
 obj, vars = is.numeric,
 tf = sqrt # SAFE: row=NULL & obs = NULL, so coercion performed
str(obj)
# Coercion examples - data.table - partial columns ====
# sb_mod():
obj <- data.table::data.table(</pre>
 a = 1:10, b = letters[1:10], c = 11:20, d = factor(letters[1:10])
str(obj) # notice that columns "a" and "c" are INTEGER (`int`)
sb2_mod(
 obj, obs = \sim (a >= 2) & (c <= 17), vars = is.numeric,
 tf = sqrt # SAFE: coercion performed
# sb_set():
obj <- data.table::data.table(</pre>
 a = 1:10, b = letters[1:10], c = 11:20, d = factor(letters[1:10])
str(obj) # notice that columns "a" and "c" are INTEGER (`int`)
 obj, obs = \sim (a >= 2) & (c <= 17), vars = is.numeric,
 tf = sqrt
 # WARNING: sqrt() results in `dbl`, but columns are `int`, so decimals lost
)
print(obj)
obj <- data.table::data.table(</pre>
 a = 1:10, b = letters[1:10], c = 11:20, d = factor(letters[1:10])
str(obj)
dt_setcoe(obj, vars = is.numeric, v = as.numeric)
str(obj)
sb2_set(obj,
 obs = \sim (a >= 2) & (c <= 17), vars = is.numeric,
 tf = sqrt # SAFE: coercion performed by dt_setcoe(); so no warnings
)
print(obj)
```

View of List ====

```
x <- list(
  a = data.table::data.table(cola = 1:10, colb = letters[1:10]),
  b = data.table::data.table(cola = 11:20, colb = letters[11:20]))
print(x)
myref <- x$a
address(myref) == address(x$a) # they are the same
sb2_set(myref, vars = "cola", tf = \(x)x^2\)
print(x) # notice x has been changed</pre>
```

aaa09_squarebrackets_slicev

On Index-Less Value-Based Sub-Set Operations

Description

This help page explains the details on the arguments used in the slicev_ methods and the county function.

The Basic Idea

The basic idea is as follows.

Let x and y be 2 atomic vectors of the same length (but they don't have to be of the same type). Let v be some atomic scalar of the same type as y.

Given the result r of the condition y == v, the basic idea is to perform the following sub-set operations:

The above is with the default argument specification r = TRUE.

Of course one can invert the relationship by specifying argument r = FALSE, to get something like the following:

And y is allowed to be the same vector as x, of course.

This basic idea, however, can become more complicated, depending on the atomic type of y, which is discussed in the next section.

Details per Atomic Type

Logical, Raw, Complex

For y of type logical, raw, and complex, slicev works exactly as explained in the previous section. y and v must be of the same atomic type.

Numeric

For y of type integer or double (collectively referred to as "numeric"), the basic idea laid-out before still holds:

one can use atomic vector y and atomic scalar v to perform sub-set operations like x[y == v].

But one may be more interested in a range of numbers, rather than one specific number (especially considering things like measurement error, and machine precision, and greater-than/larger-than relationships).

So for numeric y, one can also supply v of length 2.

When length(v) == 2L, $slicev_{red}$ countv will check whether y is inside (or outside if r = FALSE) the bounded range given by v.

I.e.

```
y \ge v[1] & y \le v[2] # if r = TRUE

y \le v[1] | y > v[2] # if r = FALSE
```

Note that y and v must both be numeric here, but they don't have to be the same type. I.e. one can have y of type integer and v of type double, without problems.

Character

For y of type character, the basic idea is still to do something like x[y == v].

When searching for string v for sub-setting purposes, one may want to take into consideration things like different spelling, spacing, or even encodings of the same string.

Implementing every form of fuzzy matching or encoding matching is computationally intensive, and also quite beyond the scope of this package.

Instead, the user may supply a character vector v of arbitrary length, containing all the variations (in terms of spelling, spacing, encoding, or whatever) of all the strings to look for.

So if a vector is given for v (instead of a single string), the following check is performed:

```
y %in% v # if r = TRUE
!y %in% v # if r = FALSE
```

Factors

Technically, a factor has the type of integer, but it has special behaviour to the extend that it is treated differently in 'R'.

It is similarly treated by the slicev_/ countv_ methods and functions.

When y is a factor, v can be given as:

- a single string (matching one of the levels of y);
- a single integer (matching one of the unique values of unclass(y));
- a factor of length 1, with the same levels and level-ordering as y.

Note that factors with NA levels are not supported, and passing such a factor to y will result in an error.

Smaller Than, Greater Than

For numeric y, one can specify a range for v, as explained earlier.

But note one can also specify something like v = c(-Inf, 4), which essentially corresponds to the condition $y \le 4$.

Thus, when v specifies a range, "greater-than" and "smaller-than" comparisons are also possible. This also holds for y of type complex.

Handling NAs and NaN

We also have to handle the NAs and NaNs.

The na argument can be used to specify what to do when a y is NA.

```
When na = FALSE, all NA values of y are always ignored.
So these are not extracted (slicev_x), replaced (slicev_set), or counted (county).
```

When na = TRUE, NA values of y are always included.

So these will be included in the extractions (slicev_x), replacements (slicev_set), and counts (county).

One can also specify na = NA, which will ignore v completely, and explicitly look for NAs/NaNs in y instead - like so:

Handling NAs works the same for all atomic types.

For y of type complex, a value y[i] is considered NA if Re(y[i]) is NA/NaN and/or Im(y[i]) is NA/NaN.

Argument v is never allowed to contain NA/NaN.

From. To

Like the slice methods, one can specify the range within to perform the sub-set operations, using the from, to arguments.

For example, if you wish to extract all values of x for which holds that y != v, but **only** want the extractions between index 10 and 100, one can specify the following:

```
slicev_x(x, y = y, v = v, r = FALSE, from = 10, to = 100.
```

Just like slice, the from, to argument can also be used for reverse the order of the result, by specifying a higher value for from than for to.

The step-size in the slicev/countv functions is always 1L (or -1L if from > to).

Also like slice, from, to can be imaginary numbers also, as explained in cp_seq.

Inverting

countv() and slicev_set() do not have an "invert" argument, and likewise there is no slicev_wo()
function.

One can only invert the sub-set condition, by specifying r = FALSE.

But r = FALSE only inverts the condition; it does not invert the range specified by from, to.

Ellipsis

The ellipsis (...) is intentionally placed right after the first argument (x in slicev_ and y in countv) to force the user to explicitly name all arguments, as doing so will avoid a lot of unnecessary confusion.

```
# basic idea ====
nms <- c(letters, LETTERS, month.abb, month.name) |> rep_len(1e6)
x <- mutable_atomic(1:1e6, names = nms)
head(x)
# memory efficient form of sum(x <= 10):
countv(x, v = c(-Inf, 10))</pre>
```

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```
# extract all elements of x with the name "a":
slicev_x(x, y = names(x), v = "a") |> head()
# find all x smaller than or equal to 5, and replace with `-1000`:
slicev\_set(x, y = x, v = c(-Inf, 5), rp = -1000L)
head(x, n = 10)
# Numeric range ====
x <- mutable_atomic(1:1e6)</pre>
head(x)
slicev_x(x, v= c(-Inf, 5)) \# x[x \le 5]
# Character ====
x <- stringi::stri_rand_shuffle(rep("hello", 1e5))</pre>
slicev_x(x, v = "hello") |> head() # find "hello"
\mbox{\tt\#} find 2 possible misspellings of "hello":
slicev_x(x, v = c("holle", "helol")) |> head()
```

bind

Dimensional Binding of Objects

Description

The bind_implementations provide dimensional binding functionalities. When possible, the bind_ functions return mutable classes.

The following implementations are available:

• bind_mat() binds dimensionless (atomic/recursive) vectors and (atomic/recursive) matrices row- or column-wise.

If the result is atomic, returns a mutable_atomic matrix; otherwise returns a recursive matrix.

- bind_array() binds (atomic/recursive) arrays and (atomic/recursive) matrices. If the result is atomic, returns a mutable_atomic array; otherwise returns a recursive array.
- bind_dt() binds data.tables and other data.frame-like objects.
 Returns a data.table.
 Faster than do.call(cbind, ...) or do.call(rbind, ...) for regular data.frame objects.

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Note that the naming convention of the binding implementations here is "bind_" followed by the **resulting class** (abbreviated).

I.e. bind_mat returns a matrix, but can bind both matrices and vectors.

And bind_array returns an array, but can bind both arrays and matrices.

And bind_dt **returns** a data.table, but can bind not only data.tables, but also most other data.frame-like objects.

Usage

```
bind_mat(arg.list, along, name_deparse = TRUE, comnames_from = 1L)
bind_array(
    arg.list,
    along,
    name_along = TRUE,
    comnames_from = 1L,
    name_flat = FALSE
)
bind_dt(arg.list, along, ...)
```

Arguments

arg.list

a list of only the appropriate objects.

If arg.list is named, its names will be used for the names of dimension along of the output, as far as possible.

along

a single integer, indicating the dimension along which to bind the dimensions.

I.e. use along = 1 for row-binding, along = 2 for column-binding, etc.

For arrays, additional flexibility is available:

- Specifying along = 0 will bind the arrays on a new dimension before the first, making along the new first dimension.
- Specifying along = n+1, with n being the last available dimension, will create an additional dimension (n+1) and bind the arrays along that new dimension

name_deparse

Boolean, for bind_mat().

Indicates if dimension along should be named. Uses the naming method from rbind/cbind itself.

comnames_from

either integer scalar or NULL, for bind_mat() and bind_array().

Indicates which object in arg.list should be used for naming the shared di-

mension.

If NULL, no communal names will be given.

For example:

When binding columns of matrices, the matrices will share the same rownames.

Using comnames_from = 10 will then result in bind_array() using rownames(arg.list[[10]])

for the rownames of the output.

name_along

Boolean, for bind_array().

Indicates if dimension along should be named.

name_flat

Boolean, for bind_array().

Indicates if flat indices should be named.

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Note that setting this to TRUE will reduce performance considerably. for performance: set to FALSE

... arguments to be passed to rbindlist.

Details

bind_array() is a modified version of the fantastic abind::abind function by Tony Plare & Richard Heiberger (2016), in the following ways:

- bind_array() primarily differs from abind::abind in that it can handle recursive arrays properly
 (the original abind::abind function would unlist everything to atomic arrays, ruining the structure).
- unlike abind: :abind, bind_array() only binds (atomic/recursive) arrays and matrices. bind_array()does not attempt to convert things to arrays when they are not arrays, but will give an error instead.

This saves computation time and prevents unexpected results.

- if bind_array() results in an atomic array, it will be a mutable_atomic array.
- bind_array() has more streamlined naming options.

bind_mat() is a modified version of rbind/cbind.

The primary differences is that bind_mat() gives an error when fractional recycling is attempted (like binding 1:3 with 1:10).

Value

The bound object.

References

Plate T, Heiberger R (2016). *abind: Combine Multidimensional Arrays*. R package version 1.4-5, https://CRAN.R-project.org/package=abind.

```
# bind_array ====

# here, atomic and recursive matrices are mixed,
# resulting in a recursive matrix

# creating the arrays
x <- c(
    lapply(1:3, \(x)sample(c(TRUE, FALSE, NA))),
    lapply(1:3, \(x)sample(1:10)),
    lapply(1:3, \(x)rnorm(10)),
    lapply(1:3, \(x)sample(letters))
)
x <- matrix(x, 4, 3, byrow = TRUE)</pre>
```

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```
dimnames(x) <- n(letters[1:4], LETTERS[1:3])</pre>
print(x)
y <- matrix(1:12, 4, 3)
print(y)
# binding the arrays
arg.list <- list(x = x, y = y)
bind_array(arg.list, along = 0L) # binds on new dimension before first
bind_array(arg.list, along = 1L) # binds on first dimension
bind_array(arg.list, along = 2L)
bind_array(arg.list, along = 3L) # bind on new dimension after last
# bind_mat ====
# here, atomic and recursive matrices are mixed,
# resulting in a recursive matrix
x <- c(
 lapply(1:3, \(x)sample(c(TRUE, FALSE, NA))),
 lapply(1:3, \(x)sample(1:10)),
 lapply(1:3, \(x)rnorm(10)),
 lapply(1:3, \(x)sample(letters))
x \leftarrow matrix(x, 4, 3, byrow = TRUE)
dimnames(x) <- n(letters[1:4], LETTERS[1:3])</pre>
print(x)
y <- matrix(1:12, 4, 3)
print(y)
bind_mat(n(x = x, y = y), 2L)
# bind_dt ====
x <- data.frame(a = 1:12, b = month.abb) # data.frame
y <- data.table::data.table(a = 1:12, b = month.abb) # data.table
bind_dt(n(x = x, y = y), 2L) # column bind
bind_dt(n(x = x, y = y), 1L) # row bind
```

Description

The mutable_atomic class is a mutable version of atomic classes.

It works exactly the same in all aspects as regular atomic classes, with only one real difference:

The 'squarebrackets' methods and functions that perform modification by reference (basically all methods and functions with "set" in the name) accept mutable_atomic, but do not accept regular atomic.

See squarebrackets_PassByReference for details.

Like data.table, [<- performs R's default copy-on-modification semantics. For modification by reference, use sb_set.

Exposed functions (beside the S3 methods):

- mutable_atomic(): create a mutable_atomic object from given data.
- couldb.mutable_atomic(): checks if an object could become mutable_atomic. An objects can become mutable_atomic if it is one of the following types: logical, integer, double, character, complex, raw. Factors can never be mutable_atomic.
- typecast.mutable_atomic() type-casts and possibly reshapes a (mutable) atomic object, and returns a mutable_atomic object.
 Does not preserve dimension names if dimensions are changed.

Usage

```
mutable_atomic(data, names = NULL, dim = NULL, dimnames = NULL)
as.mutable_atomic(x, ...)
## Default S3 method:
as.mutable_atomic(x, ...)
is.mutable_atomic(x)
couldb.mutable_atomic(x)
typecast.mutable_atomic(x, type = typeof(x), dims = dim(x))
## S3 method for class 'mutable_atomic'
c(..., use.names = TRUE)
## S3 method for class 'mutable_atomic'
x[...]
## S3 replacement method for class 'mutable_atomic'
x[...] \leftarrow value
## S3 method for class 'mutable_atomic'
format(x, ...)
## S3 method for class 'mutable_atomic'
```

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```
print(x, ...)
```

Arguments

```
data atomic vector giving data to fill the mutable_atomic object.

names, dim, dimnames

see setNames and array.

x an atomic object.

... method dependent arguments.

type a string giving the type; see typeof.

dims integer vector, giving the new dimensions.

use.names Boolean, indicating if names should be preserved.

value see Extract.
```

Value

```
For mutable_atomic(), as.mutable_atomic(), typecast.mutable_atomic():
Returns a mutable_atomic object.

For is.mutable_atomic():
Returns TRUE if the object is mutable_atomic, and returns FALSE otherwise.

For couldb.mutable_atomic():
Returns TRUE if the object is one of the following types:
logical, integer, double, character, complex, raw.
Returns FALSE otherwise.
```

Warning

Always use the exported functions given by 'squarebrackets' to create a mutable_atomic object, as they make necessary checks.

Circumventing these checks may break things!

```
x <- mutable_atomic(
   1:20, dim = c(5, 4), dimnames = list(letters[1:5], letters[1:4]))
x
typecast.mutable_atomic(x, "character")

x <- matrix(1:10, ncol = 2)
x <- as.mutable_atomic(x)
is.mutable_atomic(x)
print(x)
x[, 1]
x[] <- as.double(x)
print(x)
is.mutable_atomic(x)</pre>
```

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Construct Parameters for a Sequence Based on Margins

Description

cp_seq() returns a list of parameters to construct a sequence based on the margins of an object. It is internally used by the idx_r function and slice method.

Usage

```
cp_seq(x, m = 0L, from = NULL, to = NULL, by = 1L)
```

Arguments

x the object for which to compute margin-based sequence parameters.

m integer or complex, giving the margin(s).

For non-dimensional objects or for flat indices, specify m = 0L.

from integer or complex, of the same length as m or of length 1, specifying the from

point.

to integer or complex, of the same length as m or of length 1, specifying the maxi-

mally allowed end value.

by integer, of the same length as m or of length 1, specifying the step size.

Value

A list of the following elements:

\$start:

The actual starting point of the sequence.

This is simply from translated to regular numeric.

\$end:

The actual ending point of the sequence.

This is not the same as to.

For example, the following code:

```
seq(from = 1L, to = 10L, by = 2L)
#> [1] 1 3 5 7 9
```

specifies to = 10L.

But the sequence doesn't actually end at 10; it ends at 9.

Therefore, $cp_{seq}(x, m, 1, 10, 2)$ will return end = 9, not end = 10.

This allows the user to easily predict where an sequence given in idx_r/slice will actually end.

\$by:

This will give by, but with it's sign adjusted, if needed.

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```
$length.out:
```

The actual vector lengths the sequences would be, given the translated parameters.

Arguments Details

Multiple dimensions at once

The cp_seq function can construct the sequence parameters needed for multiple dimensions at once, by specifying a vector for m.

The lengths of the other arguments are then recycled if needed.

Using only by

If from, to are not specified, using by will construct the following sequence:

If by is positive, seq.int(1L, n, by).

If by is negative, seq.int(n, 1L, by).

Where n is the maximum index (i.e. length(x) or dim(x)[m], depending on the situation).

```
Using from, to, by
```

If from, to, by are all specified, by is stored as abs(by), and the sign of by is automatically adjusted to ensure a sensible sequence is created.

Examples

```
x <- data.frame(
    a = 1:10, b = letters[1:10], c = factor(letters[1:10]), d = -1:-10
)
print(x)
ind1 <- idx_r(x, 1, 2, 2* -1i) # rows 2:(nrow(x)-1)
sb2_x(x, ind1, 1L) # extract the row range

x <- array(1:125, c(5,5,5))
d <- 1:3
s <- idx_r(x, d, 2, 2* -1i) # 2:(n-1) for every dimension
sb_x(x, s = s, d = d) # same as x[ 2:4, 2:4, 2:4, drop = FALSE]

x <- letters
x[idx_r(x, 0, 2, 2* -1i)]</pre>
```

currentBindings

List or Lock All Currently Existing Bindings Pointing To Same Address

Description

```
currentBindings(x, action = "list")
```

lists all currently existing objects sharing the same address as x, in a given environment.

```
currentBindings(x, action = "checklock")
```

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searches all **currently existing** objects sharing the same **address** as x, in a given environment, and reports which of these are locked and which are not locked.

```
currentBindings(x, action = "lockbindings")
```

searches all **currently existing** objects sharing the same **address** as x, in a given environment, and locks them using lockBinding.

See also squarebrackets_PassByReference for information regarding the relation between locked bindings and pass-by-reference modifications.

Usage

```
currentBindings(x, action = "list", env = NULL)
```

Arguments

the existing variable whose address to use when searching for bindings.

action a single string, giving the action to perform.

Must be one of the following:

• "list" (default).

• "checklock".

• "lockbindings".

env the environment where to look for objects.

If NULL (default), the caller environment is used.

Details

The lockBinding function locks a binding of an object, preventing modification.

'R' also uses locked bindings to prevent modification of objects from package namespaces.

The pass-by-reference semantics of 'squarebrackets' in principle respect this, and disallows modification of objects by reference.

However, lockBinding does not lock the address/pointer of an object, only one particular binding of an object.

This problematic; consider the following example:

```
x <- mutable_atomic(1:16)
y <- x
lockBinding("y", environment())
sb_set(x, i = 1:6, rp = 8)</pre>
```

In the above code, x and y share the same address, thus pointing to the same memory, yet only y is actually locked.

Since x is not locked, modifying x is allowed.

But since sb_set()/sb2_set() performs modification by reference, y will still be modified, despite being locked.

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The currentBindings() function allows to user to: find all **currently existing** bindings in the **caller environment** sharing the same address as x, and locking all these bindings.

Value

```
For currentBindings(x, action = "list"):
Returns a character vector.

For currentBindings(x, action = "checklock"):
Returns a named logical vector.
The names give the names of the bindings,
and each associated value indicates whether the binding is locked (TRUE) or not locked (FALSE).

For currentBindings(x, action = "lockbindings"):
Returns VOID. It just locks the currently existing bindings.
To unlock the bindings, remove the objects (see rm).
```

Warning

The currentBindings() function only locks currently existing bindings in the specified environment:

bindings that are created **after** calling currentBindings() will not automatically be locked. Thus, every time the user creates a new binding of the same object, and the user wishes it to be locked, currentBindings() must be called again.

```
x <- as.mutable_atomic(1:10)</pre>
y <- x
lockBinding("y", environment())
currentBindings(x)
currentBindings(x, "checklock") # only y is locked
# since only y is locked, we can still modify y through x by reference:
sb_set(x, i = 1, rp = -1)
print(y) # modified!
rm(list= c("y")) # clean up
# one can fix this by locking ALL bindings:
y <- x
currentBindings(x, "lockbindings") # lock all
currentBindings(x, "checklock") # all bindings are locked, including y
# the 'squarebrackets' package respects the lock of a binding,
# provided all bindings of an address are locked;
# so this will give an error, as it should:
```

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```
if(requireNamespace("tinytest")) {
  tinytest::expect_error(
    sb_set(x, i = 1, rp = -1),
    pattern = "object is locked"
  )
}
# creating a new variable will NOT automatically be locked:
z <- y # new variable; will not be locked!</pre>
currentBindings(x, "checklock") # z is not locked
currentBindings(x, "lockbindings") # we must re-run this
currentBindings(x, "checklock") # now z is also locked
if(requireNamespace("tinytest")) {
  tinytest::expect_error( # now z is also protected
    sb_set(z, i = 1, rp = -1),
    pattern = "object is locked"
 )
}
rm(list=c("x", "y", "z")) # clean up
```

developer_ci

Construct Indices

Description

These functions construct flat or dimensional indices.

- ci_flat() constructs an integer vector flat indices.
- ci_margin() constructs an integer vector of indices for one particular dimension margin.
- ci_sub() constructs a list of integer subscripts.
- ci_df() is the same as ci_margin(), except it is specifically designed for data.frame-like objects.

It is a separate function, because things like dimnames(x)[1] and rownames(x) do not always return the same output for certain data.frame-like objects.

• ci_obs() and ci_vars() construct row and column indices, respectively, data.frame-like objects.

Usage

```
ci_flat(
    x,
    i,
    inv = FALSE,
    chkdup = FALSE,
```

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```
uniquely_named = FALSE,
  .abortcall = sys.call()
ci_margin(
  Х,
  slice,
  margin,
  inv = FALSE,
  chkdup = FALSE,
  uniquely_named = FALSE,
  .abortcall = sys.call()
)
ci_sub(
  х,
  s,
  d,
  inv = FALSE,
  chkdup = FALSE,
  uniquely_named = FALSE,
  .abortcall = sys.call()
ci_df(
  х,
  slice,
  margin,
  inv = FALSE,
  chkdup = FALSE,
  uniquely_named = TRUE,
  .abortcall = sys.call()
ci_obs(
  х,
  obs,
  inv = FALSE,
  chkdup = FALSE,
  uniquely_named = TRUE,
  .abortcall = sys.call()
)
ci_vars(
  Х,
  vars,
  inv = FALSE,
  chkdup = FALSE,
  uniquely_named = TRUE,
  .abortcall = sys.call()
```

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Arguments

```
x the object for which the indices are meant.
i, s, d, slice, margin, obs, vars, inv
See squarebrackets_indx_args.
chkdup see squarebrackets_options.
for performance: set to FALSE
uniquely_named Boolean, indicating if the user knows a-priori that the relevant names of x are unique.
If set to TRUE, speed may increase.
But specifying TRUE when the relevant names are not unique will result in incorrect output.
. abortcall environment where the error message is passed to.
```

Value

An integer vector of constructed indices.

Examples

```
x <- matrix(1:25, 5, 5)
colnames(x) <- c("a", "a", "b", "c", "d")
print(x)

bool <- sample(c(TRUE, FALSE), 5, TRUE)
int <- 1:4
chr <- c("a", "a")
cplx <- 1:4 * -1i
tci_bool(bool, nrow(x))
tci_int(int, ncol(x), inv = TRUE)
tci_chr(chr, colnames(x))
tci_cplx(cplx, nrow(x))

ci_flat(x, 1:10 * -1i)
ci_margin(x, 1:4, 2)
ci_sub(x, n(1:5 * -1i, 1:4), 1:2)</pre>
```

developer_tci

Type Cast Indices

Description

These functions typecast indices to proper integer indices.

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Usage

```
tci_bool(indx, n, inv = FALSE, .abortcall = sys.call())
tci_int(indx, n, inv = FALSE, chkdup = FALSE, .abortcall = sys.call())
tci_chr(
  indx,
  nms,
  inv = FALSE,
  chkdup = FALSE,
  uniquely_named = FALSE,
  .abortcall = sys.call()
)
tci_cplx(indx, n, inv = FALSE, chkdup = FALSE, .abortcall = sys.call())
```

Arguments

indx the indices to typecast

n the relevant size, when typecasting integer or logical indices.

Examples:

• If the target is row indices, input nrow for n.

• If the target is flat indices, input the length for n.

inv Boolean, indicating if the indices should be inverted.

See squarebrackets_indx_args.

.abortcall environment where the error message is passed to.

chkdup see squarebrackets_options.

for performance: set to FALSE

nms the relevant names, when typecasting character indices.

Examples:

- If the target is row indices, input row names for nms.
- If the target is flat indices, input flat names for nms.

uniquely_named Boolean, indicating if the user knows a-priori that the relevant names of \boldsymbol{x} are

If set to TRUE, speed may increase.

But specifying TRUE when the relevant names are not unique will result in incor-

rect output.

Value

An integer vector of type-cast indices.

```
x <- matrix(1:25, 5, 5)
colnames(x) <- c("a", "a", "b", "c", "d")
print(x)
bool <- sample(c(TRUE, FALSE), 5, TRUE)</pre>
```

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```
int <- 1:4
chr <- c("a", "a")
cplx <- 1:4 * -1i
tci_bool(bool, nrow(x))
tci_int(int, ncol(x), inv = TRUE)
tci_chr(chr, colnames(x))
tci_cplx(cplx, nrow(x))

ci_flat(x, 1:10 * -1i)
ci_margin(x, 1:4, 2)
ci_sub(x, n(1:5 * -1i, 1:4), 1:2)</pre>
```

dt

Functional Forms of data.table Operations

Description

Functional forms of special data.table operations.

These functions do not use Non-Standard Evaluation.

These functions also benefit from the security measures that 'squarebrackets' implements for the pass-by-reference semantics.

- dt_aggregate() aggregates a data.table or tidytable, and returns the aggregated copy.
- dt_setcoe() coercively transforms columns of a data.table or tidytable using pass-by-reference semantics.
- dt_setrm() removes columns of a data.table or tidytable using pass-by-reference semantics.
- dt_setadd(x, new) adds the columns from data.table/tidytable new to data.table/tidytable x, thereby modifying x using pass-by-reference semantics.
- dt_setreorder() reorders the rows and/or variables of a data. table using pass-by-reference semantics.

Usage

```
dt_aggregate(x, SDcols = NULL, f, by, order_by = FALSE)

dt_setcoe(
    x,
    vars = NULL,
    inv = FALSE,
    v,
    chkdup = getOption("squarebrackets.chkdup", FALSE)
)

dt_setrm(
    x,
    vars = NULL,
    inv = FALSE,
```

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```
chkdup = getOption("squarebrackets.chkdup", FALSE)
)

dt_setadd(x, new)

dt_setreorder(x, roworder = NULL, varorder = NULL)
```

Arguments

x a data.table or tidytable.

SDcols atomic vector, giving the columns to which the aggregation function f() is to

be applied on.

f the aggregation function

by atomic vector, giving the grouping columns.

order_by Boolean, indicating if the aggregated result should be ordered by the columns

specified in by.

vars, inv see squarebrackets_indx_args.

Duplicates are not allowed.

v the coercive transformation function

chkdup see squarebrackets_options.

for performance: set to FALSE

new a data.frame-like object.

It must have column names that do not already exist in x.

roworder a integer vector of the same length as nrow(x), giving the order in which the

rows are to be re-order. Internally, this numeric vector will be turned into an

order using order, thus ensuring it is a strict permutation of 1:nrow(x).

varorder integer or character vector of the same length as ncol(x), giving the new col-

umn order.

See data.table::setcolorder.

Details

dt_setreorder(x, roworder = roworder) internally creates a new column to reorder the data.table by, and then removes the new column.

The column name is randomized, and extra care is given to ensure it does not overwrite any existing columns.

Value

```
For dt_aggregate():
```

The aggregated data. table object.

For the rest of the functions:

Returns: VOID. These functions modify the object by reference.

Do not use assignments like $x \leftarrow dt_setcoe(x, ...)$.

Since these functions return void, you'll just get NULL.

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```
# dt_aggregate on sf-data.table ====
if(requireNamespace("sf")) {
 x <- sf::st_read(system.file("shape/nc.shp", package = "sf"))</pre>
 x <- data.table::as.data.table(x)</pre>
 x$region <- ifelse(x$CNTY_ID <= 2000, 'high', 'low')
 d.aggr <- dt_aggregate(</pre>
   x, SDcols = "geometry", f= sf::st_union, by = "region"
 head(d.aggr)
# dt_setcoe ====
obj <- data.table::data.table(a = 1:10, b = letters[1:10], c = 11:20, d = factor(letters[1:10]))
str(obj) # notice that columns "a" and "c" are INTEGER (`int`)
sb2_set(
 obj, obs = \sim (a >= 2) & (c <= 17), vars = is.numeric,
 tf = sqrt # WARNING: sqrt() results in `dbl`, but columns are `int`, so decimals lost
str(obj)
obj <- data.table::data.table(a = 1:10, b = letters[1:10], c = 11:20, d = factor(letters[1:10]))
dt_setcoe(obj, vars = is.numeric, v = as.numeric) # integers are now numeric
str(obi)
sb2_set(obj,
 obs = ^{\sim} (a >= 2) & (c <= 17), vars = is.numeric,
 tf = sqrt # SAFE: coercion performed; so no warnings
)
str(obj)
# dt_setrm ====
obj <- data.table::data.table(</pre>
 a = 1:10, b = letters[1:10], c = 11:20, d = factor(letters[1:10])
str(obj)
dt_setrm(obj, vars = 1)
str(obj)
obj <- data.table::data.table(</pre>
 a = 1:10, b = letters[1:10], c = 11:20, d = factor(letters[1:10])
str(obj)
dt_setrm(obj, vars = is.numeric)
```

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idx

Convert/Translate Indices (for Copy-On-Modify Substitution)

Description

head(obj)

head(obj)

The idx() method converts indices.

dt_setreorder(obj, roworder = n:1)

dt_setreorder(obj, varorder = ncol(obj):1)

The type of output depends on the type of input index arguments given:

- idx(x, i = i, ...) converts linear indices to a strictly positive integer vector of linear indices.
- idx(x, s = s, d = d, ...) converts dimensional indices to a strictly positive integer vector of linear indices.
- idx(x, slice = slice, margin = margin, ...) converts indices of one dimension to a strictly positive integer vector of indices for that specific dimension.

Vectors (both atomic and recursive) only have index argument i.

Data.frame-like objects only have the slice, margin argument pair.

Arrays (both atomic and recursive) have the s, d argument pair, as well as the i argument and the slice, margin argument pair.

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The result of the idx() method can be used inside the regular square-brackets operators. For example like so:

```
x <- array(...)
my_sub2ind <- idx(x, s, d)
x[my_sub2ind] <- value

y <- data.frame(...)
rows <- idx(y, 1:10, 1, inv = TRUE)
cols <- idx(y, c("a", "b"), 2)
y[rows, cols] <- value</pre>
```

thus allowing the user to benefit from the convenient index translations from 'squarebrackets', whilst still using R's default copy-on-modification semantics (instead of the semantics provided by 'squarebrackets').

Usage

```
idx(x, ...)
## Default S3 method:
idx(x, i, inv = FALSE, ..., chkdup = getOption("squarebrackets.chkdup", FALSE))
## S3 method for class 'array'
idx(
  х,
  s = NULL,
  d = 1:ndim(x),
  slice = NULL,
  margin = NULL,
  i = NULL,
  inv = FALSE,
  chkdup = getOption("squarebrackets.chkdup", FALSE)
## S3 method for class 'data.frame'
idx(
  Χ,
  slice,
  margin,
  inv = FALSE,
  chkdup = getOption("squarebrackets.chkdup", FALSE)
```

Arguments

x vector, matrix, array, or data.frame; both atomic and recursive objects are supported.

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```
... see squarebrackets_method_dispatch.
i, s, d, margin, slice, inv
See squarebrackets_indx_args.
Duplicates are not allowed.
chkdup see squarebrackets_options.
for performance: set to FALSE
```

Value

```
For idx(x, i = i, ...) and idx(x, s = s, d = d, ...):
A strictly positive integer vector of flat indices.

For idx(x, margin = margin, slice = slice, ...):
A strictly positive integer vector of indices for the dimension specified in margin.
```

```
print(x)

x <- data.frame(
    a = sample(c(TRUE, FALSE, NA), 10, TRUE),
    b = 1:10,
    c = rnorm(10),
    d = letters[1:10],
    e = factor(letters[11:20])
)

rows <- idx(x, 1:5, 1, inv = TRUE)
cols <- idx(x, c("b", "a"), 2)
x[rows, cols] <- NA</pre>
```

 idx_by

```
print(x)
```

idx_by

Compute Grouped Indices

Description

Given:

- a sub-set function f;
- an object x with its margin m;
- and a grouping factor grp;

the idx_by() function takes indices per group grp.

The result of idx_by() can be supplied to the indexing arguments (see squarebrackets_indx_args) to perform **grouped** subset operations.

Usage

```
idx_by(x, m, f, grp, parallel = FALSE, mc.cores = 1L)
```

Arguments

the object from which to compute the indices.

a single non-negative integer giving the margin for which to compute indices.
For flat indices or for non-dimensional objects, use m = 0L.

a subset function to be applied per group on indices.
If m == 0L, indices is here defined as setNames(1:length(x), names(x)).
If m > 0L, indices is here defined as setNames(1:dim(x)[m], dimnames(x)[[m]]).
The function must produce a character or integer vector as output.
For example, to subset the last element per group, specify:
f = last

grp a factor giving the groups.

parallel, mc.cores

Value

A vector of indices.

see BY.

 idx_ord_v 61

Examples

```
# vectors ====
(a <- 1:20)
(grp <- factor(rep(letters[1:5], each = 4)))</pre>
# get the last element of `a` for each group in `grp`:
s <- list(idx_by(a, 0L, last, grp))</pre>
sb_x(cbind(a, grp), s, 1L)
# data.frame ====
x <- data.frame(</pre>
 a = sample(1:20),
 b = letters[1:20],
 group = factor(rep(letters[1:5], each = 4))
print(x)
# get the first row for each group in data.frame `x`:
row <- idx_by(x, 1, first, x$group)</pre>
sb2_x(x, row, 1L)
\# get the first row for each group for which a > 10:
x2 <- sb2_x(x, obs = ~a > 10)
row <- na.omit(idx_by(x2, 1, first, x2$group))</pre>
sb2_x(x2, row, 1L)
```

idx_ord_v

Compute Ordered Indices

Description

Computes ordered indices. Similar to order, except the user must supply a vector, a list of equallength vectors, a data.frame or a matrix (row-wise and column-wise are both supported), as the input.

```
For a vector x, idx_ord_v(x) is equivalent to order(x).

For a data.frame or a list of equal-length vectors x, with p columns/elements, idx_ord_df(x) is equivalent to order(x[[1]], ..., x[[p]]).

For a matrix (or array) x with p rows, idx_ord_m(x, margin = 1) is equivalent to order(x[1, ], ..., x[p, ], ...).

For a matrix (or array) x with p columns, idx_ord_m(x, margin = 2) is equivalent to order(x[, 1], ..., x[, p], ...).
```

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Note that these are merely convenience functions, and that these are actually slightly slower than order (except for idx_ord_v()), due to the additional functionality.

Usage

```
idx_ord_v(
 х,
 na.last = TRUE,
 decr = FALSE,
 method = c("auto", "shell", "radix")
idx_ord_m(
 Х,
 margin,
 na.last = TRUE,
 decr = FALSE,
 method = c("auto", "shell", "radix")
idx_ord_df(
 Х,
 na.last = TRUE,
 decr = FALSE,
 method = c("auto", "shell", "radix")
```

Arguments

Value

See order.

```
x <- sample(1:10)
order(x)
idx_ord_v(x)
idx_ord_m(rbind(x, x), 1)
idx_ord_m(cbind(x, x), 2)
idx_ord_df(data.frame(x, x))</pre>
```

 idx_r 63

idx_r

Compute Integer Index Range

Description

```
idx_r() computes integer index range(s).
```

Usage

```
idx_r(x, m = 0L, from = NULL, to = NULL, by = 1L)
```

Arguments

```
x the object for which to compute subset indices. 
m, from, to, by see cp_seq.
```

Value

```
If length(m) == 1L: a vector of numeric indices.
```

If length(m) > 1L: a list of the same length as m, containing numeric vectors of indices.

```
x <- data.frame(
    a = 1:10, b = letters[1:10], c = factor(letters[1:10]), d = -1:-10
)
print(x)
ind1 <- idx_r(x, 1, 2, 2* -1i) # rows 2:(nrow(x)-1)
sb2_x(x, ind1, 1L) # extract the row range

x <- array(1:125, c(5,5,5))
d <- 1:3
s <- idx_r(x, d, 2, 2* -1i) # 2:(n-1) for every dimension
sb_x(x, s = s, d = d) # same as x[ 2:4, 2:4, 2:4, drop = FALSE]

x <- letters
x[idx_r(x, 0, 2, 2* -1i)]</pre>
```

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indx_x

Exported Utilities

Description

Exported utilities.

Usually the user won't need these functions.

Usage

```
indx_x(i, x, xnames, xsize)
indx_wo(i, x, xnames, xsize)
```

Arguments

i See squarebrackets_indx_args.

x a vector, vector-like object, factor, data.frame, data.frame-like object, or a list.

xnames names or dimension names xsize length or dimension size

Value

The subsetted object.

Examples

```
x <- 1:10
names(x) <- letters[1:10]
indx_x(1:5, x, names(x), length(x))
indx_wo(1:5, x, names(x), length(x))</pre>
```

lst

Unnest Tree-like List into a Recursive Matrix or Flattened Recursive Vector

Description

[[, [[<-, sb2_rec, and sb2_recin, can perform recursive subset operations on a nested list. Such recursive subset operations only operate on a single element.

Performing recursive subset operations on multiple elements is not vectorized, and requires a (potentially slow) loop.

The lst_untree() function takes a nested tree-like list, and turns it into a recursive matrix (a matrix of list-elements), allowing vectorized subset operations to be performed on the nested list.

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lst_untree() can also simply flatten the list, making it a non-nested list. See the Examples section to understand how the list will be arranged and named.

The lst_nlists() counts the total number of recursive list-elements inside a list.

Usage

```
lst_nlists(x)

lst_untree(x, margin, use.names = TRUE)
```

Arguments

x a tree-like nested list.

margin a single integer, indicating how the result should be arranged:

- margin = 0 produces a simple flattened recursive vector (i.e. list) without dimensions.
- margin = 1 produces a recursive matrix (i.e. a matrix of list-elements), with length(x) rows and n columns, where n = sapply(x, lst_nlists) |> max().
 Empty elements will be filled with list(NULL).
- margin = 2 produces a recursive matrix (i.e. a matrix of list-elements), with length(x) columns and n rows, where n = sapply(x, lst_nlists) |> max().
 Empty elements will be filled with list(NULL).

use.names

Boolean, indicating if the result should be named. See section "use.names" for more information.

Value

```
For lst_untree():
```

A non-nested (dimensional) list.

Note that if margin = 1 or margin = 2, lst_untree() returns a recursive matrix (i.e. a recursive array with 2 dimensions), **not** a data.frame.

To turn a nested list into a data.frame instead, one option would be to use:

```
rrapply(x, how = "melt")
```

For lst_nlists():

A single integer, giving the total number of recursive list-elements in the given list.

use.names

```
margin = 0 and use.names = TRUE
```

If margin = 0 and use.names = TRUE, every element in the flattened list will be named.

Names of nested elements, such as x[["A"]][["B"]][["C"]], will become "A.B.C", as that is the behaviour of the rapply function (which lst_untree() calls internally).

It is therefore advised not to use dots (".") in your list names, and use underscores ("_") instead, before calling lst_untree().

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See the rrapply::rrapply function for renaming (and other forms of transforming) recursive subsets of lists.

```
margin = 1 and use.names = TRUE
```

If margin == 1 and use.names = TRUE, the rows of resulting recursive matrix will be equal to names(x), but recursive names will not be assigned.

```
margin = 2 and use.names = TRUE
```

If margin == 2 and use.names = TRUE, the columns of resulting recursive matrix will be equal to names(x), but recursive names will not be assigned.

```
use.names = FALSE
```

If use.names = FALSE, the result will not have any names assigned at all.

```
# show-casing how the list-elements are arranged and named ====
x <- list(
  A = list(
   A = list(A = "AAA", B = "AAB"),
    A = list(A = "AA2A", B = "AA2B"),
   B = list(A = "ABA", B = "ABB"),
   C = letters
  ),
  Y = list(
   Z = list(Z = "YZZ", Y = "YZY"),
   Y = list(Z = "YYZ", Y = "YYY"),
   X = "YX"
 )
)
# un-tree column-wise:
sapply(x, lst_nlists) |> max() # number of rows `y` will have
y <- lst_untree(x, margin = 2L, use.names = TRUE)</pre>
dim(y)
print(y)
sb2_x(y, n(1:3, 1:2), 1:ndim(y)) # vectorized selection of multiple recursive elements
# un-tree row-wise:
sapply(x, lst_nlists) |> max() # number of columns `y` will have
y <- lst_untree(x, margin = 1L, use.names = TRUE)</pre>
dim(y)
print(y)
sb2_x(y, n(1:2, 1:3), 1:ndim(y)) # vectorized selection of multiple recursive elements
# simple flattened list:
y <- lst_untree(x, margin = 0, use.names = TRUE)</pre>
print(y)
y[["Y.Z.Y"]]
x[[c("Y", "Z", "Y")]] # equivalent in the original list
```

match_all 67

```
# showcasing that only list-elements are recursively flattened ====
# i.e. atomic vectors in recursive subsets remain atomic
x <- lapply(1:10, \(x)list(sample(letters), sample(1:10)))</pre>
sapply(x, lst_nlists) |> max()
y <- lst_untree(x, margin = 1)</pre>
dim(y)
print(y)
lst\_untree(x, margin = 1)
# showcasing vectorized sub-setting ====
x \leftarrow lapply(1:10, \(x) list(
 list(sample(letters[1:10]), sample(LETTERS[1:10])),
 list(sample(month.abb), sample(month.name)),
 list(sample(1:10), rnorm(10))
))
y \leftarrow lst\_untree(x, 1)
# getting the first recursive elements in the second level/depth in base R:
for(i in seq_along(x)) {
 x[[c(i, c(1L, 1L))]] \mid > print() \# for-loop, slow
# the same, but vectorized using the untree'd list:
sb2_x(y, n(1:nrow(y), 1L), 1:ndim(y)) > drop() > print() # vectorized, fast
```

match_all

Match All, Order-Sensitive and Duplicates-Sensitive

Description

Find all indices of vector haystack that are equal to vector needles, taking into account the order of both vectors, and their duplicate values.

match_all() is essentially a much more efficient version of:

```
lapply(needles, \(i) which(haystack == i))
```

Like lapply(needles, (i) which(haystack == i)), NAs are ignored.

match_all() internally calls collapse::fmatch and collapse::gsplit.

Core of the code is based on a suggestion by Sebastian Kranz (author of the 'collapse' package).

68 n

Usage

```
match_all(needles, haystack, unlist = TRUE)
```

Arguments

needles, haystack

vectors of the same type.

needles cannot contain NA/NaN. Long vectors are not supported.

unlist

Boolean, indicating if the result should be a single unnamed integer vector (TRUE, default), or a named list of integer vectors (FALSE).

Value

An integer vector, or list of integer vectors.

If a list, each element of the list corresponds to each value of needles.

When needles and/or haystack is empty, or when haystack is fully NA, match_all() returns an empty integer vector (if unlist = TRUE), or an empty list (if unlist = FALSE).

Examples

```
n <- 200
haystack <- sample(letters, n, TRUE)
needles <- sample(letters, n/2, TRUE)
indices1 <- match_all(needles, haystack)
head(indices1)</pre>
```

n Nest

Description

The c() function concatenates vectors or lists into a vector (if possible) or else a list.

In analogy to that function, the n() function **nests** objects into a list (not into an atomic vector, as atomic vectors cannot be nested).

It is a short-hand version of the list function.

This is handy because lists are often needed in 'squarebrackets', especially for arrays.

Usage

n()

Value

The list.

ndim 69

Examples

```
obj <- array(1:64, c(4,4,3))
print(obj)
sb_x(obj, n(1:3, 1:2), c(1,3))
# above is equivalent to obj[1:3, , 1:2, drop = FALSE]</pre>
```

ndim

Get Number of Dimensions

Description

```
ndim(x) is short-hand for length(dim(x)).
```

Usage

ndim(x)

Arguments

Х

the object to get the number of dimensions from.

Value

An integer, giving the number of dimensions x has. For vectors, gives $\emptyset L$.

```
x <- 1:10
ndim(x)
obj <- array(1:64, c(4,4,3))
print(obj)
ndim(obj)</pre>
```

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sb2_rec

Access, Replace, Transform, Delete, or Extend Recursive Subsets

Description

The sb2_rec() and sb2_recin() methods are essentially convenient wrappers around [[and [[<-, respectively.

Unlike [[and [[<-, these are actually S3 methods, so package authors can create additional method dispatches.

sb2_rec() will access recursive subsets of lists.

sb2_recin() can do the following things:

- replace or transform recursive subsets of a list, using R's default Copy-On-Modify semantics, by specifying the rp or tf argument, respectively.
- delete a recursive subset of a list, using R's default Copy-On-Modify semantics, by specifying argument rp = NULL.
- extending a list with additional recursive elements, using R's default Copy-On-Modify semantics

This is done by specifying an out-of-bounds index in argument rec, and entering the new values in argument rp.

Note that adding surface level elements of a dimensional list will delete the dimension attributes of that list.

Usage

```
sb2_rec(x, ...)
## Default S3 method:
sb2_rec(x, rec, ...)
sb2_recin(x, ...)
## Default S3 method:
sb2_recin(x, rec, ..., rp, tf)
```

Arguments

x a list, or list-like object.

... see squarebrackets_method_dispatch.

rec a strictly positive integer vector or character vector, of length p, such that sb2_rec(x, rec) is equivalent to x[[rec[1]]]...[[rec[p]]], providing all but the final indexing results in a list.

When on a certain subset level of a nested list, multiple subsets with the same name exist, only the first one will be selected when performing recursive indexing by name, since recursive indexing can only select a single element.

NA, NaN, Inf, -Inf are not valid values for rec.

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rp optional, and allows for multiple functionalities:

• In the simplest case, performs x[[rec]] <- rp, using R's default semantics. Since this is a replacement of a recursive subset, rp does not necessarily have to be a list itself; rp can be any type of object.

- Specifying rp = NULL will **delete** (recursive) subset sb(x, rec).

 To specify actual NULL instead of deleting a subset, use rp = list(NULL).
- When rec is an integer, and specifies an out-of-bounds subset, sb2_recin() will add value rp to the list.

Any empty positions in between will be filled with NA.

• When rec is character, and specifies a non-existing name, sb2_recin() will add value rp to the list as a new element at the end.

tf

an optional function. If specified, performs $x[[rec]] \leftarrow tf(x[[rec]])$, using R's default Copy-On-Modify semantics.

Does not support extending a list like argument rp.

Details

Since recursive objects are references to other objects, extending a list or deleting an element of a list does not copy the entire list, in contrast to atomic vectors.

Value

```
For sb2_rec():
```

Returns the recursive subset.

```
For sb2\_recin(..., rp = rp):
```

Returns VOID, but replaces, adds, or deletes the specified recursive subset, using R's default Copy-On-Modify semantics.

```
For sb2\_recin(..., tf = tf):
```

Returns VOID, but transforms the specified recursive subset, using R's default Copy-On-Modify semantics.

```
lst <- list(
    A = list(
    A = list(A = "AAA", B = "AAB"),
    A = list(A = "AA2A", B = "AA2B"),
    B = list(A = "ABA", B = "ABB")
),
    B = list(
    A = list(A = "BAA", B = "BAB"),
    B = list(A = "BBA", B = "BBB")
),
    C = list(
    A = 1:10,</pre>
```

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```
B = 11:20
# access recursive subsets ====
sb2_rec(lst, c(1,2,2)) # this gives "AA2B"
sb2_rec(lst, c("A", "B", "B")) # this gives "ABB"
sb2_rec(lst, c(2,2,1)) # this gives "BBA"
sb2_rec(lst, c("B", "B", "A")) # this gives "BBA"
# replace recursive subset with R's default in-place semantics ====
# replace "AAB" using R's default in-place semantics:
sb2_recin(
 lst, c("A", "A", "B"),
 rp = "THIS IS REPLACED WITH IN-PLACE SEMANTICS"
print(lst)
# transform recursive subsets with R's default in-place semantics ====
sb2\_recin(lst, c("C", "A"), tf = \(x)x^2) # transforms lst$C$A
print(lst)
# add/remove new recursive subsets with R's default in-place semantics ====
sb2_recin(lst, c("C", "D"), rp = "NEW VALUE") # adds lst$C$D
print(lst)
sb2_recin(lst, c("C", "A"), rp = NULL) # removes lst$C$A
print(lst) # notice lst$C$A is GONE
# Modify View of List By Reference ====
x <- list(
a = data.table::data.table(cola = 1:10, colb = letters[1:10]),
b = data.table::data.table(cola = 11:20, colb = letters[11:20])
)
print(x)
myref <- sb2_rec(x, "a")
```

```
address(myref) == address(xa) # they are the same sb2_set(myref, vars = "cola", tf = \(x)x^2\) print(x) # notice x has been changed
```

 sb_mod

Method to Return a Copy of an Object With Modified Subsets

Description

This is an S3 Method to return a copy of an object with modified subsets. Use $sb_mod(x, ...)$ if x is an atomic object; this returns a full copy. Use $sb2_mod(x, ...)$ if x is a recursive object (i.e. list or data.frame-like); this returns a partial copy.

For modifying subsets using R's default copy-on-modification semantics, see idx.

```
sb_mod(x, ...)
## Default S3 method:
sb_mod(
  Х,
  i = NULL,
  inv = FALSE,
  ...,
  rp,
  tf,
  chkdup = getOption("squarebrackets.chkdup", FALSE)
## S3 method for class 'array'
sb_mod(
  Х,
  s = NULL,
  d = 1:ndim(x),
  i = NULL,
  inv = FALSE,
  . . . ,
  rp,
  tf,
  chkdup = getOption("squarebrackets.chkdup", FALSE)
sb2\_mod(x, ...)
## Default S3 method:
```

```
sb2_mod(
  Х,
  i = NULL,
  inv = FALSE,
  ...,
  rp,
  tf,
  chkdup = getOption("squarebrackets.chkdup", FALSE),
  .lapply = lapply
## S3 method for class 'array'
sb2_mod(
  х,
  s = NULL,
  d = 1:ndim(x),
  i = NULL,
  inv = FALSE,
  rp,
  tf,
  chkdup = getOption("squarebrackets.chkdup", FALSE),
  .lapply = lapply
## S3 method for class 'data.frame'
sb2_mod(
  Х,
  s = NULL,
  d = 1:2,
  obs = NULL,
  vars = NULL,
  inv = FALSE,
  rp,
  tf,
  chkdup = getOption("squarebrackets.chkdup", FALSE),
  .lapply = lapply
```

Arguments

```
x see squarebrackets_supported_structures.
... see squarebrackets_method_dispatch.
i, s, d, obs, vars, inv
See squarebrackets_indx_args.
An empty index selection returns the original object unchanged.
rp, tf, .lapply see squarebrackets_modify.
chkdup see squarebrackets_options.
for performance: set to FALSE
```

Details

Transform or Replace

Specifying argument tf will transform the subset.

Specifying rp will replace the subset.

One cannot specify both tf and rp. It's either one set or the other.

Value

A copy of the object with replaced/transformed values.

```
# atomic objects ====
obj \leftarrow matrix(1:16, ncol = 4)
colnames(obj) \leftarrow c("a", "b", "c", "a")
print(obj)
rp <- -1:-9
sb_mod(obj, n(1:3), 1:ndim(obj), rp = rp)
# above is equivalent to obj[1:3, 1:3] <- -1:-9; obj</pre>
sb_{mod}(obj, i = (x)x \le 5, rp = -1:-5)
# above is equivalent to obj[obj <= 5] <- -1:-5; obj</pre>
sb_{mod}(obj, n("a"), 2L, rp = -1:-8)
# above is equivalent to obj[, which(colnames(obj) %in% "a")] <- -1:-8; obj
sb_{mod(obj, n(1:3), 1:ndim(obj), tf = (x) -x)
# above is equivalent to obj[1:3, 1:3] \leftarrow (-1 * obj[1:3, 1:3]); obj
sb_mod(obj, i = \(x)x \le 5, tf = \(x) -x)
# above is equivalent to obj[obj \le 5] <- (-1 * obj[obj \le 5]); obj
obj <- array(1:64, c(4,4,3))
print(obj)
sb_{mod}(obj, n(1:3, 1:2), c(1,3), rp = -1:-24)
# above is equivalent to obj[1:3, , 1:2] <- -1:-24
sb_{mod}(obj, i = (x)x \le 5, rp = -1:-5)
# above is equivalent to obj[obj <= 5] <- -1:-5
# lists ====
obj <- list(a = 1:10, b = letters[1:11], c = 11:20)
print(obj)
sb2_mod(obj, "a", rp = list(1L))
# above is equivalent to obj[["a"]] <- 1L; obj</pre>
sb2_{mod}(obj, is.numeric, rp = list(-1:-10, -11:-20))
# above is equivalent to obj[which(sapply(obj, is.numeric))] <- list(-1:-10, -11:-20); obj</pre>
obj <- rbind(</pre>
  lapply(1:4, \(x)sample(c(TRUE, FALSE, NA))),
  lapply(1:4, \(x)sample(1:10)),
  lapply(1:4, \xspace (x)rnorm(10)),
```

```
lapply(1:4, \(x)sample(letters))
colnames(obj) <- c("a", "b", "c", "a")</pre>
print(obj)
sb2_mod(obj, n(1:3), 1:ndim(obj), rp = n(-1))
# above is equivalent to obj[1:3, 1:3] <- list(-1)</pre>
sb2_mod(obj, i = is.numeric, rp = n(-1))
# above is equivalent to obj[sapply(obj, is.numeric)] <- list(-1)</pre>
sb2_{mod}(obj, n("a"), 2L, rp = n(-1))
# above is equivalent to
\# obj[, lapply(c("a", "a"), \(i) which(colnames(obj) == i)) |> unlist()] <- list(-1)
obj <- array(as.list(1:64), c(4,4,3))
print(obj)
sb2_mod(obj, n(1:3, 1:2), c(1,3), rp = as.list(-1:-24))
\# above is equivalent to obj[1:3, , 1:2] <- as.list(-1:-24)
sb2_mod(obj, i = \(x) x \le 5, rp = as.list(-1:-5))
# above is equivalent to obj[sapply(onj, (x) \times (5)] <- as.list(-1:-5)
# data.frame-like objects - whole columns ====
obj <- \ data.frame(a = 1:10, \ b = letters[1:10], \ c = 11:20, \ d = factor(letters[1:10]))\\
str(obj) # notice that columns "a" and "c" are INTEGER (`int`)
sb2_mod(
 obj, vars = is.numeric,
 tf = sqrt
# data.frame-like objects - partial columns ====
obj <- data.frame(a = 1:10, b = letters[1:10], c = 11:20, d = factor(letters[1:10]))
str(obj) # notice that columns "a" and "c" are INTEGER (`int`)
sb2 mod(
 obj, obs = \sim (a >= 2) & (c <= 17), vars = is.numeric,
 tf = sqrt
sb2_mod(
 obj, obs = \sim (a >= 2) & (c <= 17), vars = is.numeric,
  tf = sqrt
sb2_mod(
 obj, obs = \sim (a >= 2) & (c <= 17), vars = is.numeric,
 tf = sqrt
```

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sb_set

Method to Modify Subsets of a Mutable Object By Reference

Description

This is an S3 Method to replace or transform a subset of a supported mutable object using pass-by-reference semantics

```
Use sb\_set(x, ...) if x is an atomic object (i.e. mutable_atomic). Use sb2\_set(x, ...) if x is a recursive object (i.e. data.table).
```

```
sb_set(x, ...)
## Default S3 method:
sb_set(
  Х,
  i = NULL,
  inv = FALSE,
  . . . ,
  rp,
  tf,
  chkdup = getOption("squarebrackets.chkdup", FALSE)
)
## S3 method for class 'array'
sb_set(
  Х,
  s = NULL,
  d = 1:ndim(x),
  i = NULL,
  inv = FALSE,
  . . . ,
  rp,
  tf,
  chkdup = getOption("squarebrackets.chkdup", FALSE)
sb2\_set(x, ...)
## Default S3 method:
sb2\_set(x, ...)
## S3 method for class 'data.table'
sb2_set(
  х,
  s = NULL,
  d = 1:2,
  obs = NULL,
```

```
vars = NULL,
inv = FALSE,
...,
rp,
tf,
chkdup = getOption("squarebrackets.chkdup", FALSE),
.lapply = lapply
)
```

Arguments

```
x a variable belonging to one of the supported mutable classes.

... see squarebrackets_method_dispatch.
i, s, d, obs, vars, inv
See squarebrackets_indx_args.
An empty index selection leaves the original object unchanged.

rp, tf, .lapply see squarebrackets_modify.
chkdup see squarebrackets_options.
for performance: set to FALSE
```

Details

Transform or Replace

Specifying argument tf will transform the subset. Specifying rp will replace the subset. One cannot specify both tf and rp. It's either one set or the other.

Value

```
Returns: VOID. This method modifies the object by reference. Do not use assignments like x <- sb_set(x, ...). Since this function returns void, you'll just get NULL.
```

```
# mutable_atomic objects ====
gen_mat <- function() {
  obj <- as.mutable_atomic(matrix(1:16, ncol = 4))
  colnames(obj) <- c("a", "b", "c", "a")
  return(obj)
}
obj <- obj2 <- gen_mat()
print(obj)</pre>
```

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```
sb_set(obj, n(1:3), 1:ndim(obj), rp = -1:-9)
print(obj2)
# above is like x[1:3, 1:3] <- -1:-9, but using pass-by-reference
obj <- obj2 <- gen_mat()</pre>
obj
sb_set(obj, i = \(x) x \le 5, rp = -1:-5)
print(obj2)
# above is like x[x \le 5] < -1:-5, but using pass-by-reference
obj <- obj2 <- gen_mat()</pre>
sb_set(obj, n("a"), 2L, rp = cbind(-1:-4, -5:-8))
print(obj2)
# above is like x[, "a"] <- cbind(-1:-4, -5:-8), but using pass-by-reference
obj <- obj2 <- gen_mat()</pre>
obj
sb_set(obj, n(1:3), 1:ndim(obj), tf = \(x) -x)
# above is like x[1:3, 1:3] \leftarrow -1 * x[1:3, 1:3], but using pass-by-reference
obj <- obj2 <- gen_mat()</pre>
obj
sb_set(obj, i = \(x) x \le 5, tf = \(x) -x)
print(obj2)
# above is like x[x \le 5] < -1 * x[x \le 5], but using pass-by-reference
obj <- obj2 <- gen_mat()</pre>
obj
sb_set(obj, n("a"), 2L, tf = \(x) -x)
obj2
# above is like x[, "a"] <- -1 * x[, "a"], but using pass-by-reference
gen_array <- function() {</pre>
 as.mutable_atomic(array(1:64, c(4,4,3)))
obj <- obj2 <- gen_array()</pre>
obj
sb_set(obj, n(1:3, 1:2, c(1, 3)), 1:3, rp = -1:-12)
print(obj2)
# above is like x[1:3, , 1:2] \leftarrow -1:-12, but using pass-by-reference
obj <- obj2 <- gen_array()
obj
sb_set(obj, i = (x)x \le 5, rp = -1:-5)
print(obj2)
# above is like x[x \le 5] < -1:-5, but using pass-by-reference
```

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```
# data.table ====
obj \leftarrow data.table::data.table(a = 1:10, b = letters[1:10], c = 11:20, d = factor(letters[1:10]))
str(obj) # notice that columns "a" and "c" are INTEGER (`int`)
sb2 set(
  obj, obs = \sim (a >= 2) & (c <= 17), vars = is.numeric,
  tf = sqrt # WARNING: sqrt() results in `dbl`, but columns are `int`, so decimals lost
print(obj)
obj <- data.table::data.table(a = 1:10, b = letters[1:10], c = 11:20, d = factor(letters[1:10]))
dt_setcoe(obj, vars = is.numeric, v = as.numeric)
str(obj)
sb2_set(obj,
 obs = ^{\sim} (a >= 2) & (c <= 17), vars = is.numeric,
  tf = sqrt # SAFE: coercion performed by dt_setcoe(); so no warnings
print(obj)
obj <- data.table::data.table(a = 1:10, b = letters[1:10], c = 11:20, d = factor(letters[1:10]))
str(obj) # notice that columns "a" and "c" are INTEGER (`int`)
sb2_set(
 obj, vars = is.numeric,
  tf = sqrt # SAFE: row=NULL & obs = NULL, so coercion performed
str(obj)
```

sb_setRename

Safely Change the Names of a Mutable Object By Reference

Description

Functions to rename a supported mutable object using pass-by-reference semantics:

- \bullet sb_setFlatnames() renames the (flat) names of a mutable_atomic object.
- sb_setDimnames() renames the dimension names of a mutable_atomic object.
- \bullet sb2_setVarnames() renames the variable names of a data.table object.

```
sb_setFlatnames(x, i = NULL, newnames, ...)
sb_setDimnames(x, m, newdimnames, ...)
sb2_setVarnames(x, old, new, skip_absent = FALSE, ...)
```

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Arguments x

i logical, numeric, character, or imaginary indices, indicating which flatnames

a variable belonging to one of the supported mutable classes.

should be changed.

If i = NULL, the names will be completely replaced.

newnames Atomic character vector giving the new names.

Specifying NULL will remove the names.

... see squarebrackets_method_dispatch.

m integer vector giving the margin(s) for which to change the names (m = 1L for

rows, m = 2L for columns, etc.).

newdimnames a list of the same length as m.

The first element of the list corresponds to margin m[1], the second element to

m[2], and so on.

The components of the list can be either NULL, or a character vector with the

same length as the corresponding dimension.

Instead of a list, simply NULL can be specified, which will remove the dimnames

completely.

old the old column names

mutable atomic matrix ====

new the new column names, in the same order as old

 $skip_absent$ Skip items in old that are missing (i.e. absent) in names(x).

Default FALSE halts with error if any are missing.

Value

Returns: VOID. This method modifies the object by reference. Do not use assignment like names(x) <- sb_setRename(x, ...). Since this function returns void, you'll just get NULL.

sb_wo

sb_wo

print(x)

Method to Return Object Without Specified Subset

Description

```
This is an S3 Method to return an object without the specified subset. sb_wo()/sb_wo() is essentially the inverse of sb_x/sb_x. Use sb_wo(x, ...) if x is an atomic object. Use sb_wo(x, ...) if x is a recursive object (i.e. list or data.frame-like).
```

```
sb_wo(x, ...)
## Default S3 method:
sb_wo(x, i = NULL, ..., chkdup = getOption("squarebrackets.chkdup", FALSE))
## S3 method for class 'array'
sb_wo(
    x,
    s = NULL,
    d = 1:ndim(x),
    i = NULL,
```

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. . . ,

red

```
chkdup = getOption("squarebrackets.chkdup", FALSE)
   sb2\_wo(x, ...)
   ## Default S3 method:
    sb2_wo(
      х,
      i = NULL,
      red = FALSE,
      chkdup = getOption("squarebrackets.chkdup", FALSE)
   ## S3 method for class 'array'
    sb2_wo(
      х,
      s = NULL,
      d = 1:ndim(x),
      i = NULL,
      red = FALSE,
      chkdup = getOption("squarebrackets.chkdup", FALSE)
   ## S3 method for class 'data.frame'
    sb2_wo(
      х,
      s = NULL,
      d = 1:2,
      obs = NULL,
      vars = NULL,
      chkdup = getOption("squarebrackets.chkdup", FALSE)
Arguments
                    see squarebrackets_supported_structures.
    Χ
                    see squarebrackets_method_dispatch.
    i, s, d, obs, vars See squarebrackets_indx_args.
                    An empty index selection results in nothing being removed, and the entire object
                    is returned.
                    see squarebrackets_options.
   chkdup
```

for performance: set to FALSE

ing [[]].

Boolean, for recursive objects only, indicating if the result should be reduced.

If red = TRUE, selecting a single element will give the simplified result, like us-

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If red = FALSE, a list is always returned regardless of the number of elements.

Value

A copy of the sub-setted object.

Examples

```
# atomic objects ====
obj <- matrix(1:16, ncol = 4)
colnames(obj) <- c("a", "b", "c", "a")</pre>
print(obj)
sb_wo(obj, n(1:3), 1:ndim(obj))
# above is equivalent to obj[-1:-3, -1:-3, drop = FALSE]
sb_wo(obj, i = (x) x > 5)
# above is equivalent to obj[!obj > 5]
sb_wo(obj, n("a"), 2L)
# above is equivalent to obj[, which(!colnames(obj) %in% "a")]
obj <- array(1:64, c(4,4,3))
print(obj)
sb_{wo}(obj, n(1, c(1, 3)), c(1, 3))
# above is equivalent to obj[-1, , c(-1, -3), drop = FALSE]
sb_wo(obj, i = (x)x > 5)
# above is equivalent to obj[!obj > 5]
```

```
# lists ====
obj <- list(a = 1:10, b = letters[1:11], c = 11:20)
print(obj)
sb2_wo(obj, "a")
# above is equivalent to obj[which(!names(obj) %in% "a")]
sb2_wo(obj, 1) # obj[-1]
sb2_wo(obj, 1:2)
# above is equivalent to obj[seq_len(length(obj))[-1:-2]]
sb2_wo(obj, is.numeric, red = TRUE)
# above is equivalent to obj[[!sapply(obj, is.numeric)]] IF this returns a single element
obj <- list(a = 1:10, b = letters[1:11], c = letters)
sb2_wo(obj, is.numeric)
# above is equivalent to obj[!sapply(obj, is.numeric)] # this time singular brackets?
# for recusive indexing, see sb2_rec()
obj <- rbind(</pre>
  lapply(1:4, \(x)sample(c(TRUE, FALSE, NA))),
  lapply(1:4, \(x)sample(1:10)),
  lapply(1:4, \(x)rnorm(10)),
 lapply(1:4, \(x)sample(letters))
colnames(obj) <- c("a", "b", "c", "a")</pre>
```

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```
print(obj)
sb2_wo(obj, n(1:3), 1:ndim(obj))
# above is equivalent to obj[1:3, 1:3, drop = FALSE]
sb2\_wo(obj, i = is.numeric)
# above is equivalent to obj[sapply(obj, is.numeric)]
sb2_wo(obj, n(c("a", "a")), 2L)
# above is equivalent to obj[, lapply(c("a", "a"), (i) which(colnames(obj) == i)) |> unlist()]
obj <- array(as.list(1:64), c(4,4,3))
print(obj)
sb2_wo(obj, n(1, c(1, 3)), c(1, 3))
# above is equivalent to obj[-1, , c(-1, -3), drop = FALSE]
sb2\_wo(obj, i = \(x)x>5)
# above is equivalent to obj[!sapply(obj, (x) x > 5)]
# data.frame-like objects ====
obj <- data.frame(a = 1:10, b = letters[1:10], c = 11:20, d = factor(letters[1:10]))
print(obj)
sb2_wo(obj, n(1:3))
# above is equivalent to obj[-1:-3, -1:-3, drop = FALSE]
sb2\_wo(obj, obs = ~(a > 5) & (c < 19), vars = is.numeric)
```

sb_x

Method to Extract, Exchange, or Duplicate Subsets of an Object

Description

```
This is an S3 Method to extract, exchange, or duplicate (i.e. repeat x times) subsets of an object. Use sb_x(x, ...) if x is an atomic object. Use sb_x(x, ...) if x is a recursive object (i.e. list or data.frame-like).
```

```
sb_x(x, ...)
## Default S3 method:
sb_x(x, i = NULL, ...)
## S3 method for class 'array'
sb_x(x, s = NULL, d = 1:ndim(x), i = NULL, ...)
sb2_x(x, ...)
```

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```
## Default S3 method:
sb2_x(x, i = NULL, red = FALSE, ...)
## S3 method for class 'array'
sb2_x(x, s = NULL, d = 1:ndim(x), i = NULL, red = FALSE, ...)
## S3 method for class 'data.frame'
sb2_x(x, s = NULL, d = 1:2, obs = NULL, vars = NULL, ...)
```

Arguments

x see squarebrackets_supported_structures.
... see squarebrackets_method_dispatch.
i, s, d, obs, vars See squarebrackets_indx_args.
Duplicates are allowed, resulting in duplicated indices.
An empty index selection results in an empty object of length 0.

red Boolean, for recursive objects only, indicating if the result should be reduced.

If red = TRUE, selecting a single element will give the simplified result, like using [[]].

If red = FALSE, a list is always returned regardless of the number of elements.

Value

Returns a copy of the sub-setted object.

Examples

```
# atomic objects ====
obj \leftarrow matrix(1:16, ncol = 4)
\texttt{colnames(obj)} \; \mathrel{<\!\!\!\!-} \; \mathsf{c("a", "b", "c", "a")}
print(obj)
sb_x(obj, s = n(1:3), d = 1:ndim(obj))
# above is equivalent to obj[1:3, 1:3, drop = FALSE]
sb_x(obj, i = (x) x > 5)
# above is equivalent to obj[obj > 5]
sb_x(obj, s = n(c("a", "a")), d = 2L)
\# above is equivalent to obj[, lapply(c("a", "a"), \(i) which(colnames(obj) == i)) |> unlist()]
obj <- array(1:64, c(4,4,3))
print(obj)
sb_x(obj, s = n(1:3, 1:2), d = c(1,3))
# above is equivalent to obj[1:3, , 1:2, drop = FALSE]
sb_x(obj, i = (x)x > 5)
# above is equivalent to obj[obj > 5]
```

```
# lists ====
```

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```
obj <- list(a = 1:10, b = letters[1:11], c = 11:20)
print(obj)
sb2_x(obj, 1) # obj[1]
sb2_x(obj, 1, red = TRUE) # obj[[1]]
sb2_x(obj, 1:2) # obj[1:2]
sb2_x(obj, is.numeric) # obj[sapply(obj, is.numeric)]
# for recursive subsets, see sb2_rec()
obj <- rbind(</pre>
  lapply(1:4, \(x)sample(c(TRUE, FALSE, NA))),
  lapply(1:4, \x) sample(1:10)),
  lapply(1:4, \x)rnorm(10)),
 lapply(1:4, \(x)sample(letters))
colnames(obj) <- c("a", "b", "c", "a")</pre>
print(obj)
sb2_x(obj, s = n(1:3), d = 1:ndim(obj))
# above is equivalent to obj[1:3, 1:3, drop = FALSE]
sb2_x(obj, i = is.numeric)
# above is equivalent to obj[sapply(obj, is.numeric)]
sb2_x(obj, s = n(c("a", "a")), d = 2L)
# above is equivalent to obj[, lapply(c("a", "a"), \(i) which(colnames(obj) == i)) |> unlist()]
obj <- array(as.list(1:64), c(4,4,3))
print(obj)
sb2_x(obj, s = n(1:3, 1:2), d = c(1,3))
# above is equivalent to obj[1:3, , 1:2, drop = FALSE]
sb2_x(obj, i = (x)x > 5)
# above is equivalent to obj[sapply(obj, (x) \times 5)]
# data.frame-like objects ====
obj <- data.frame(a = 1:10, b = letters[1:10], c = 11:20, d = factor(letters[1:10]))
print(obj)
sb2_x(obj, n(1:3)) # obj[1:3, 1:3, drop = FALSE]
sb2_x(obj, obs = ~(a > 5) & (c < 19), vars = is.numeric)
```

setapply

Apply Functions Over mutable_atomic Matrix Margins By Reference

Description

The setapply() function applies a functions over the rows or columns of a mutable_atomic matrix, through pass-by-reference semantics.

The setapply() is a bit faster and uses less memory than apply.

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Usage

```
setapply(x, MARGIN, FUN)
```

Arguments

x a mutable_atomic 2-dimensional array (i.e. a matrix).

Arrays of other than 2 dimensions are not supported.

MARGIN a single integer scalar, giving the subscript to apply the function over.

1 indicates rows, 2 indicates columns.

FUN the function to be applied.

The function must return a vector of the same type of x, and the appropriate length (i.e. length ncol(x) when MARGIN == 1 or length nrow(x) when MARGIN

== 2).

Value

Returns: VOID. This function modifies the object by reference. Do NOT use assignment like x <- setapply(x, ...). Since this function returns void, you'll just get NULL.

Examples

```
# re-order elements matrix by reference ====
x <- mutable_atomic(1:20, dim = c(5,4))
print(x)
setapply(x, 1, FUN = \(x)x[c(4,1,3,2)])
print(x)

# sort elements of matrix by reference ====
x <- mutable_atomic(20:1, dim = c(5,4))
print(x)
setapply(x, 2, FUN = sort)
print(x)</pre>
```

slice

Efficient Sequence-based Subset Methods on (Long) Vectors

Description

The slice_ - methods are similar to the sb_ - methods, except they don't require an indexing vector, and are designed for memory efficiency.

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Usage

```
slice_x(x, ...)
## Default S3 method:
slice_x(
 х,
 from = NULL,
 to = NULL,
 by = 1L,
 use.names = TRUE,
 sticky = getOption("squarebrackets.sticky", FALSE)
slice_wo(x, ...)
## Default S3 method:
slice_wo(
 х,
 from = NULL,
 to = NULL,
 by = 1L,
 ...,
 use.names = TRUE,
 sticky = getOption("squarebrackets.sticky", FALSE)
slice_set(x, ...)
## Default S3 method:
slice_set(x, from = NULL, to = NULL, by = 1L, inv = FALSE, ..., rp, tf)
```

Arguments

х	an atomic object. For slice_set it must be a mutable_atomic variable.
	see squarebrackets_method_dispatch.
from, to, by	see cp_seq.
use.names	Boolean, indicating if flat names should be preserved. Note that, since the slice_ methods operates on flat indices only, dimensions and dimnames are always dropped.
sticky	see squarebrackets_options.
inv	Boolean, indicating whether to invert the sequence. If TRUE, slice_set() will apply replacement/transformation on all elements of the vector, except for the elements of the specified sequence.
rp, tf	see squarebrackets_modify.

Value

Similar to the sb_ methods.

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Examples

```
x <- mutable_atomic(1:1e7)

# extract:
slice_x(x, 1, 10)

# reverse:
slice_x(x, -1i, 1) |> head()

# remove:
slice_wo(x, 1, -11i) # all elements except the last 10

# replace every other element:
x <- mutable_atomic(1:1e7)
slice_set(x, 2, -1i, 2, rp = -1)
head(x)

# replace all elements except the first element:
x <- mutable_atomic(1:1e7)
slice_set(x, 1, 1, inv = TRUE, rp = -1)
head(x)</pre>
```

slicev

Efficient Value-based Subset Methods on (Long) Vectors

Description

The slicev_ - methods are similar to the sb_ - methods, except they don't require an indexing vector, and are designed for memory efficiency.

counv(y, v, from, to) counts how often a value, or range of values, v, occurs in a vector subset y[from:to].

```
slicev_x(x, ...)
## Default S3 method:
slicev_x(
    x,
    ...,
    y = x,
    v = NULL,
    na = FALSE,
    r = TRUE,
    from = NULL,
    to = NULL,
    use.names = TRUE,
```

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```
sticky = getOption("squarebrackets.sticky", FALSE)
)
slicev_set(x, ...)

## Default S3 method:
slicev_set(
    x,
    ...,
    y = x,
    v = NULL,
    na = FALSE,
    r = TRUE,
    from = NULL,
    to = NULL,
    rp,
    tf
)
countv(y, ..., v = NULL, na = FALSE, r = TRUE, from = NULL, to = NULL)
```

Arguments

X	an atomic vector. For slicev_set() it must be a mutable_atomic variable.
	See squarebrackets_slicev.
y, v, na, r	See squarebrackets_slicev.
from, to	see cp_seq.
use.names	Boolean, indicating if flat names should be preserved. Note that, since the slicev_ methods operates on flat indices only, dimensions and dimnames are always dropped.
sticky	see squarebrackets_options.
rp, tf	see squarebrackets_modify.

Value

Similar to the sb_ methods.

For countv(): A single number, giving the number of elements matching the specified condition.

```
# basic idea ====
nms <- c(letters, LETTERS, month.abb, month.name) |> rep_len(1e6)
x <- mutable_atomic(1:1e6, names = nms)
head(x)

# memory efficient form of sum(x <= 10):
countv(x, v = c(-Inf, 10))</pre>
```

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```
# extract all elements of x with the name "a":
slicev_x(x, y = names(x), v = "a") |> head()
# find all x smaller than or equal to 5, and replace with `-1000`:
slicev_set(x, y = x, v = c(-Inf, 5), rp = -1000L)
head(x, n = 10)
# Numeric range ====
x <- mutable_atomic(1:1e6)</pre>
head(x)
slicev_x(x, v= c(-Inf, 5)) # x[x <= 5]
# Character ====
x <- stringi::stri_rand_shuffle(rep("hello", 1e5))</pre>
head(x)
slicev_x(x, v = "hello") |> head() # find "hello"
# find 2 possible misspellings of "hello":
slicev_x(x, v = c("holle", "helol")) |> head()
```

sub2ind

Convert Subscripts to Coordinates, Coordinates to Flat Indices, and Vice-Versa

Description

These functions convert a list of integer subscripts to an integer matrix of coordinates, an integer matrix of coordinates to an integer vector of flat indices, and vice-versa. Inspired by the sub2ind function from 'MatLab'.

- sub2coord() converts a list of integer subscripts to an integer matrix of coordinates.
- coord2ind() converts an integer matrix of coordinates to an integer vector of flat indices.
- ind2coord() converts an integer vector of flat indices to an integer matrix of coordinates.
- coord2sub() converts an integer matrix of coordinates to a list of integer subscripts; it performs a very simple (one might even say naive) conversion.
- sub2ind() is a faster and more memory efficient version of coord2ind(sub2coord(sub, x.dims), x.dims)

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All of these functions are written to be memory-efficient.

The coord2ind() is thus the opposite of arrayInd, and ind2coord is merely a convenient wrapper around arrayInd.

Note that the equivalent to the sub2ind function from 'MatLab' is actually the coord2ind() function here.

Usage

```
sub2coord(sub, x.dim)
coord2sub(coord)
coord2ind(coord, x.dim, checks = TRUE)
ind2coord(ind, x.dim)
sub2ind(sub, x.dim, checks = TRUE)
```

Arguments

sub a list of integer subscripts.

The first element of the list corresponds to the first dimension (rows), the second

element to the second dimensions (columns), etc.

The length of sub must be equal to the length of x.dim.

One cannot give an empty subscript; instead fill in something like seq_len(dim(x)[margin]).

NOTE: The coord2sub() function does not support duplicate subscripts.

x. dim an integer vector giving the dimensions of the array in question. I.e. dim(x).

coord an integer matrix, giving the coordinate indices (subscripts) to convert.

Each row is an index, and each column is the dimension.

The first columns corresponds to the first dimension, the second column to the

second dimensions, etc.

The number of columns of coord must be equal to the length of x. dim.

checks Boolean, indicating if arguments checks should be performed.

Defaults to TRUE.

Can be set to FALSE for minor speed improvements.

for performance: set to FALSE

ind an integer vector, giving the flat position indices to convert.

Details

The base S3 vector and array classes in 'R' use the standard Linear Algebraic convention, as in academic fields like Mathematics and Statistics, in the following sense:

- vectors are **column** vectors (i.e. vertically aligned vectors);
- index counting starts at 1;
- rows are the first dimension/subscript, columns are the second dimension/subscript, etc.

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Thus, the orientation of flat indices in, for example, a 4-rows-by-5-columns matrix, is as follows:

```
[,1] [,2] [,3] [,4] [,5]
[1,]
        1
             5
                  9
                      13
[2,]
        2
             6
                 10
                       14
                            18
[3,]
        3
             7
                 11
                       15
                            19
[4,]
                 12
                       16
                            20
```

So in a 4 by 5 matrix, subscript [1, 2] corresponds to flat index 5. Array subscripting in 'squarebrackets' also follows this standard convention.

Value

For sub2coord() and ind2coord():

Returns an integer matrix of coordinates (with properties as described in argument coord).

For coord2ind():

Returns an numeric vector of flat indices (with properties as described in argument ind).

For coord2sub():

Returns a list of integer subscripts (with properties as described in argument sub)

For sub2ind():

Returns an integer vector of flat indices(if $prod(x.dim) < (2^31 - 1)$), or an numeric vector of flat indices (if $prod(x.dim) >= (2^31 - 1)$).

Note

These functions were not specifically designed for duplicate indices per-sé. For efficiency, they do not check for duplicate indices either.

```
x.dim <- c(10, 10, 3)
x.len <- prod(x.dim)
x <- array(1:x.len, x.dim)
sub <- list(c(4, 3), c(3, 2), c(2, 3))
coord <- sub2coord(sub, x.dim)
print(coord)
ind <- coord2ind(coord, x.dim)
print(ind)
all(x[ind] == c(x[c(4, 3), c(3, 2), c(2, 3)])) # TRUE
coord2 <- ind2coord(ind, x.dim)
print(coord)
all(coord == coord2) # TRUE
sub2 <- coord2sub(coord2)
sapply(1:3, \(i) sub2[[i]] == sub[[i]]) |> all() # TRUE
```

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