The \textit{scidb} Package

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The scidb Package

1 Introduction

SciDB is an open-source database that organizes data in \( n \)-dimensional arrays. SciDB features include ACID transactions, parallel processing, distributed storage, efficient sparse array storage, and native linear algebra operations. The scidb package for R provides two ways to interact with SciDB from R:

1. Through SciDB array classes for R. The arrays mimic standard R arrays and data frames, but operations on them are performed by the SciDB engine. Data are materialized to R only when requested.
2. By running SciDB queries from R, optionally transferring data through data frames or data frame iterators.

The SciDB array classes facilitate programming large-scale SciDB computation in R using natural R syntax. This vignette illustrates using SciDB from R by example. For more detailed information on the functions described in this vignette, see the manual pages in the package.

2 Connecting to SciDB

The scidbconnect function establishes a connection to a simple HTTP network service called shim running on a SciDB coordinator instance (see Section 12.3). The function may be safely called multiple times. Once a connection is established, connection information is maintained until a different connection is established or the R session ends.

The network interface optionally supports SSL encryption and user authentication. Users are defined by the operating system users on the SciDB coordinator instance. The shim service can be configured to support either open/unencrypted or encrypted/authenticated ports, or both.

Connect to localhost by default on unencrypted port 8080:

```
> library("scidb")
> scidbconnect()
```

Connect to SciDB on an encrypted port 8083 with example authentication:

```
> scidbconnect(host="localhost", port=8083, 
+ username="scidbuser", password="test")
```
We recommend using only encrypted/authenticated sessions when communicating with SciDB over public networks.

2.1 Listing and removing SciDB arrays

The `scidblist` function lists SciDB objects (arrays, instances, etc.), optionally showing detailed schema information for arrays. Returned results may be filtered using regular expression-style syntax.

The `scidbremove` function removes a SciDB array, or optionally a set of SciDB arrays defined by regular expression. The function accepts a vector of array names, resulting in the removal of all the specified arrays. Combine this feature with the regular expression filtering output of `scidblist` to remove sets of arrays matching the filter.

3 SciDB Arrays for R users

Data are organized by SciDB in \(n\)-dimensional sparse arrays. "Sparse" in SciDB arrays means that array elements may be left undefined, and such array elements are omitted from computations. Note that this interpretation of sparse differs in a subtle way from that used by sparse matrices defined by R’s Matrix package (whose sparse matrix elements are implicitly zero).

The elements of a SciDB array, called *cells*, contain one or more *attributes* (similar to R variables). The number and data types of attributes are uniform across all cells in an array, and SciDB stores data for each attribute separately. Thus, a one-dimensional SciDB array is conceptually very similar to a data frame in R: the SciDB dimension index corresponds to data frame row index, and SciDB attributes to data frame columns. Higher-dimensional arrays in SciDB don’t correspond directly to objects in R; the `scidb` \(n\)-dimensional array class described below is limited to working with one attribute at a time to more closely emulate standard R arrays.

The `scidb` package defines two array classes for R with data backed by SciDB arrays: the `scidbdf` class for data frame-like objects, and the `scidb` class for matrix and general array objects.

3.1 The `scidbdf` data frame-like class

The `scidbdf` class defines a data frame-like class backed by one-dimensional SciDB arrays. Like data frames, the columns represent variables of distinct types and the rows represent observations.
Each attribute in the backing SciDB array represents a column in the \texttt{scidbdf} object. The \texttt{scidbdf} object elements are read-only (the backing SciDB array may be manually updated, for example using the \texttt{iquery} function).

Use the \texttt{as.scidb} function to create new SciDB arrays and corresponding \texttt{scidbdf} R objects by copying R data frames into SciDB. The \texttt{types} and \texttt{nullable} options may be used to explicitly specify the SciDB type and nullability values of each data frame column. See the R help page for \texttt{as.scidb} and related \texttt{df2scidb} functions for more information.

The \texttt{scidb} function returns an R \texttt{scidbdf} or \texttt{scidb} object representation of an existing SciDB array.

Objects of class \texttt{scidbdf} obey a subset of R indexing operations. Columns may be selected by numeric positional index, variable name, or with the shorthand dollar sign notation similarly to standard data frames. Rows may be selected by numeric ranges—see the following section on array indexing for details.

Subsets of \texttt{scidbdf} objects are returned as new \texttt{scidbdf} objects of the appropriate size (dimension, number of attributes/columns). The package uses the special empty-bracket notation, [], to indicate that data should be materialized to R as an R data frame. Illustrations are provided in the examples.

```r
> # Upload the Michelson-Morley experiment data to SciDB,
> # returning a \texttt{scidbdf} data frame-like object:
> X <- as.scidb(morley, name="morely")
> str(X)

SciDB expression: morely
SciDB schema: <Expt:int32 NULL DEFAULT null,Run:int32 NULL DEFAULT null,Speed:int32 NULL DEFAULT null>[row=1:100,100,0]

Attributes:
  attribute type nullable
  1   Expt  int32   TRUE
  2    Run  int32   TRUE
  3   Speed  int32   TRUE

Dimension:
  dimension start end chunk
  1    row   1 100   100

> # Materialize the first four rows of \texttt{X} and the "Run" and "Speed"
```
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> # columns to R (using [] to return results to R):
> X[1:4, c("Run","Speed")][]

<table>
<thead>
<tr>
<th>Run</th>
<th>Speed</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>850</td>
</tr>
<tr>
<td>2</td>
<td>740</td>
</tr>
<tr>
<td>3</td>
<td>900</td>
</tr>
<tr>
<td>4</td>
<td>1070</td>
</tr>
</tbody>
</table>

3.2 The `scidb` array class

Similarly to the data frame-like class, the package defines the `scidb` array class for R that represents vectors, matrices and general n-dimensional arrays. Array objects defined by the `scidb` class behave in some ways like standard R arrays. But their data reside in SciDB and most operations on them are computed by SciDB.

The `scidb` array class typically support working with a single array attribute at a time to conform to R arrays (which generally support a single value per cell). Note that, unlike R, SciDB array origin indices are arbitrary and may be zero or negative (matrices and vectors in SciDB are generally zero-indexed). Data from `scidb` array objects are not materialized to R until extracted with the empty indexing function,[]. Additional notes and examples about indexing appear in the next section.

> # Upload the iris data to SciDB storing it in an array named `iris`
> # and create a data frame-like object `df` that refers to it:
> df <- as.scidb(iris)
> dim(df)

[1] 150 5

> # Alternatively, make a SciDB 1-d array object.
> x <- scidb(df, data.frame=FALSE)
> dim(x)

[1] 150

> # List all available attributes in the SciDB array:
> x@attributes
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[1] "Sepal_Length" "Sepal_Width" "Petal_Length" "Petal_Width" "Species"

3.3 Arithmetic operations

The scidb array class supports a few standard linear algebra operations for dense and sparse matrices and vectors.

The example below shows a dense matrix example that compares matrix arithmetic in R and SciDB. Like previous data frame examples, use `as.scidb` to export R matrices and vectors to SciDB arrays.

We generally use `as.scidb` for convenience—it’s far from the most efficient way to import data into SciDB. For very large data, use the SciDB bulk data load utility as outlined in the SciDB documentation instead.

```r
> v <- as.scidb(matrix(rnorm(25),5))
> str(v)

SciDB expression: R_array2bb317c6889d1101074813220
SciDB schema: <val:double NULL DEFAULT null> [i=0:4,5,0,j=0:4,5,0]

Attributes:

<table>
<thead>
<tr>
<th>attribute type</th>
<th>nullable</th>
</tr>
</thead>
<tbody>
<tr>
<td>val double</td>
<td>TRUE</td>
</tr>
</tbody>
</table>

Dimensions:

<table>
<thead>
<tr>
<th>dimension start end chunk</th>
</tr>
</thead>
<tbody>
<tr>
<td>i 0 4 5</td>
</tr>
<tr>
<td>j 0 4 5</td>
</tr>
</tbody>
</table>

> crossprod( v[] )  # Compute t(v) %*% v using R

```

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>9.532370</td>
<td>-7.472959</td>
<td>4.952529</td>
<td>3.736609</td>
<td>1.464429</td>
</tr>
<tr>
<td>-7.472959</td>
<td>8.079816</td>
<td>-2.546614</td>
<td>-2.024801</td>
<td>-2.133998</td>
</tr>
<tr>
<td>4.952529</td>
<td>-2.546614</td>
<td>3.895251</td>
<td>2.868848</td>
<td>1.625864</td>
</tr>
<tr>
<td>3.736609</td>
<td>-2.024801</td>
<td>2.868848</td>
<td>2.18813</td>
<td>1.237344</td>
</tr>
<tr>
<td>1.464429</td>
<td>-2.133998</td>
<td>1.625864</td>
<td>1.237344</td>
<td>5.295838</td>
</tr>
</tbody>
</table>
```
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> # Now compute using SciDB, and materialize the result to R:
> `crossprod(v)[]`

```
[1,]  9.532370 -7.472959  4.952529  3.736609  1.464429
[3,]  4.952529 -2.546614  3.895251  2.868848  1.625864
[4,]  3.736609 -2.024801  2.868848  2.218813  1.237344
[5,]  1.464429 -2.133998  1.625864  1.237344  5.295838
```

Basic matrix/vector arithmetic operations on SciDB arrays (addition, subtraction, matrix and matrix vector products, scalar products, `crossprod` and `tcrossprod`) use standard R syntax. You can mix R and SciDB matrices and vectors and the `scidb` package will try to do the right thing by assigning R data to temporary SciDB arrays conforming to required database schema. The next example shows an example of computations that mix `scidb` array objects with R vectors.

<table>
<thead>
<tr>
<th>Expression</th>
<th>Operation</th>
<th>Operands</th>
<th>Output</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>A %*% B</code></td>
<td>Matrix multiplication</td>
<td><code>A,B</code> Conformable SciDB arrays or R matrices/vectors</td>
<td>SciDB array</td>
</tr>
<tr>
<td><code>A ± B</code></td>
<td>Matrix summation/difference</td>
<td><code>A,B</code> SciDB arrays or R matrices/vectors</td>
<td>SciDB array</td>
</tr>
<tr>
<td><code>crossprod(A,B)</code></td>
<td>Cross product</td>
<td><code>A,B</code> SciDB arrays or R matrices/vectors</td>
<td>SciDB array</td>
</tr>
<tr>
<td><code>tcrossprod(A,B)</code></td>
<td>Cross product</td>
<td><code>A,B</code> SciDB arrays or R matrices/vectors</td>
<td>SciDB array</td>
</tr>
<tr>
<td><code>A */ B</code></td>
<td>Elementwise product/quotient</td>
<td><code>A,B</code> Conformable SciDB arrays or R matrices/vectors</td>
<td>SciDB array</td>
</tr>
<tr>
<td><code>α */ A</code></td>
<td>Scalar multiplication/division</td>
<td>SciDB array A, scalar <code>α</code></td>
<td>SciDB array</td>
</tr>
<tr>
<td><code>t(A)</code></td>
<td>Transpose</td>
<td>SciDB array A</td>
<td>SciDB array</td>
</tr>
<tr>
<td><code>sin(A)</code></td>
<td>Sine function, also other trig functions</td>
<td>SciDB array A</td>
<td>SciDB array</td>
</tr>
<tr>
<td><code>log(A, base)</code></td>
<td>Logarithm function</td>
<td>SciDB array A, numeric base</td>
<td>SciDB array</td>
</tr>
<tr>
<td><code>diff(A, lag = 1)</code></td>
<td>Finite differences</td>
<td>SciDB array A, integer lag</td>
<td>SciDB array</td>
</tr>
<tr>
<td><code>A[range, range,...]</code></td>
<td>Subarray</td>
<td>SciDB array A</td>
<td>SciDB Array</td>
</tr>
<tr>
<td><code>A[]</code></td>
<td>Materialize</td>
<td>SciDB array A</td>
<td>R array</td>
</tr>
<tr>
<td><code>diag(A)</code></td>
<td>Matrix diagonal</td>
<td>SciDB array or vector A</td>
<td>SciDB arrays</td>
</tr>
<tr>
<td><code>svd(A)</code></td>
<td>Singular value decompostion</td>
<td>Dense SciDB array A</td>
<td>SciDB arrays</td>
</tr>
<tr>
<td><code>svd(A, nu)</code></td>
<td>Truncated SVD</td>
<td>Sparse or dense SciDB array A</td>
<td>SciDB arrays</td>
</tr>
</tbody>
</table>
4 Subsetting and indexing SciDB objects

The `scidb` and `scidbdf` classes generally follow SciDB database indexing convention, which exhibits some differences with standard R indexing. In particular, note that the starting SciDB integer index is arbitrary, but often zero. The upper-left corner of R arrays is always indexed by \([1,1,...]\). Subarray indexing operations use the SciDB convention. Thus, zero and negative indices are literally interpreted and passed to SciDB. In particular, negative indices do not indicate index omission, unlike standard R arrays.

Important indexing notes include:

- Use empty brackets, [], to materialize data back to R. Otherwise, indexing operations produce new SciDB array objects.

- Use numeric indices in any dimension in the units of the underlying SciDB array coordinate system. Note that SciDB arrays generally are zero-indexed and may even have negative indices. SciDB data frame-like objects generally use 1-based indexing. When in doubt about the base index, use the `str` or `dimensions` functions to interrogate the SciDB object for details.

- Numeric indexing may include contiguous ranges or vectors of distinct coordinate values, but repeated coordinate values in a single dimension are not allowed right now (sorry about that). Examples of valid index ranges include \([1:4, \ c(3,1,5),\ -10:15]\), but not \([c(1,3,1)]\).

- The `scidbdf` class represents 1-d SciDB arrays as data frame objects with SciDB array attribute as columns. Use either positional numeric or name-based indexing along columns, either with the dollar-sign notation or string indexing.

- The `scidb` supports labeled dimension indexing using R `rownames`, `colnames`, or `dimnames` settings. Labels assigned in this way must be provided by 1-d SciDB arrays that map the integer coordinates to character label values. See the examples below.

- The `scidb` class supports indexing by other SciDB arrays to achieve the effect of filtering by boolean expressions and similar operations, also illustrated below in the examples.

- Use the utility `between` function to avoid forming large sequences representing huge indexing ranges. For example, use `[between(1,1e9)]` instead of `[1:1e9]`.

- The `diag` function is supported to extract a vector of diagonal elements from a matrix, or to create a sparse diagonal matrix from a vector.
4.1 Comparison operators and filtering

The package follows R convention and returns a logical-valued object after comparison. Additionally, the package defines special comparison operators enclosed in the `%` symbol that return a sparse SciDB array object whose values not meeting the condition are masked as empty (preserving the values that do meet the condition).

The difference is illustrated in the following example.

```r
> x <- as.scidb(head(iris))
> # Standard R-style logical comparison:
> (x$Petal_Length > 1.4)[]

[1] FALSE FALSE FALSE TRUE FALSE TRUE

> # Special SciDB-style masked comparison:
> (x$Petal_Length %>% 1.4)[]

sparse vector (nnz/length = 2/6) of class "dsparseVector"
[1] . . . 1.5 . 1.7
```

Comparison results can be used as filters in scidb and scidbdf objects. This works mostly like R, except that such filters may only be applied along coordinate axes. (That is, R’s special single-index indexing mode is not supported.)

Additional examples showing dimension labels and filtering follow.

```r
> # Example of labeling indices with a label array.
> set.seed(1)
> X <- as.scidb( matrix(rnorm(20),nrow=5) )
> # SciDB matrix objects like X default to zero-based indexing.
> # It's important that the label array have the same starting index:
> rownames(X) <- as.scidb( data.frame(letters[1:5]), start=0)
> X[c("b","a","d"), ]

A reference to a 3x4 SciDB array
```

```r
> # Example of indexing by a condition on an auxiliary array.
> # Determine a subset of the rownames of X (another SciDB array):
> idx <- rownames(X) > "b"
> # Subset X by the condition on its rows:
> X[idx, ]
```
5 Data manipulation functions

The package defines a number of common SciDB data manipulation functions for SciDB array objects.

5.1 Apply, sweep and other array functions

R’s apply function applies a function along margins of an array or matrix. A version of apply limited to SciDB aggregation functions is available for scidb and scidbdf objects. The SciDB aggregation function is supplied as a string and references the array attributes. The next example uses apply together with sweep to center the columns of a matrix.

```r
> # Create an example matrix and upload to SciDB. Its single numeric array
> # attribute will be called "val" be default.
> A <- as.scidb(matrix(rnorm(100), nrow=10))
> # Create a centered version of the matrix by computing its column means
> # with apply and subtracting them with sweep. Note that apply uses a
> # SciDB expression. We specify `eval=TRUE` below to compute and cache
> # the result of sweep, improving the efficiency of the subsequent
> # crossprod operation.
> A0 <- sweep(A, 2, apply(A, 2, "avg(val)"), eval=TRUE)
> # The covariance matrix of A:
> crossprod(A0)/(nrow(A) - 1)
```

A reference to a 10x10 SciDB array

More general aggregation and data manipulation functions are outlined in the next section. The diff function works similarly to R’s usual one for computing finite differences along vectors and matrices.

Use the cumulate function to compute running operations along data, for example cumulative sums. The operation to be performed must be a valid SciDB aggregation function expressed as a character string. Here is a simple example:
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```r
> x <- as.scidb(iris)
> y <- cumulate(x, "sum(Petal_Width)"
> print(head(y, n=4))

[1] 0.2 0.4 0.6 0.8
```

The `count` function applied to a `scidb` array object returns the count of non-empty cells in the backing SciDB array.

The `image` function displays a heatmap of a regrid of a 2-d `scidb` array object, and returns the regridded array to R. The `grid=c(m,n)` function parameter specifies the regrid window sizes in each array dimension, and defaults to the array chunk sizes. The regrid aggregation function may be specified using the `op` function argument, and by default averages the array values over the regrid windows.

### 5.2 Conditional subset operations

Use the `subset` function to filter array contents by a boolean expression somewhat similarly to the standard R `subset` function. Under the hood, this function uses the SciDB `filter` operator—the function name `subset` more closely matches standard R syntax.

The `subset` function requires two arguments, a SciDB array reference and a valid SciDB logical expression represented as a string. Here is a simple example:

```r
> df <- as.scidb(iris)
> subset(df,"Petal_Width > 2.4")

    Sepal_Length Sepal_Width Petal_Length Petal_Width Species
   101       6.3        3.3         6.0  2.5 virginica
   110       7.2        3.6         6.1  2.5 virginica
   145       6.7        3.3         5.7  2.5 virginica
```

The `subset` function applied to arrays returns a sparse SciDB array with entries not meeting the filter condition omitted. This behavior differs from the standard R `subset` function, which can only operate on rows and columns. The `scidb` class also supports a short-hand notation for `subset` for infix comparison operations.

```r
> # Upload a 5x5 matrix to SciDB. Its single numeric attribute name 
> # defaults to 'val' (see help for as.scidb):
```
> set.seed(1)
> A <- as.scidb( matrix(rnorm(9),nrow=3) )
> # Apply a filter to the SciDB array and return the results to R:
> subset(A, "val>0")[]

3 x 3 sparse Matrix of class "dgCMatrix"

[1,] . 1.5952808 0.4874291
[2,] 0.1836433 0.3295078 0.7383247
[3,] . . 0.5757814

> # Alternatively, achieve the same filter with an infix comparison:
> (A > 0)[]

[,1] [,2] [,3]
[1,] FALSE TRUE TRUE
[2,] TRUE TRUE TRUE
[3,] FALSE FALSE TRUE

### 5.3 Database joins

The package supports a limited form of the R `merge` function, enabling a number of database join-like operations on SciDB array objects. See `help("merge",package="scidb")` function for detailed help. Here is an example that performs an inner join on array attributes:

> authors <-
+ data.frame(  
+   surname = c("Tukey", "Venables", "Tierney", "Ripley", "McNeil"),
+   nationality = c("US", "Australia", "US", "UK", "Australia"),
+   deceased = c("yes", rep("no", 4)),
+   stringsAsFactors=FALSE)
> books <-
+ data.frame(  
+   name = c("Tukey", "Venables", "Tierney",
+   "Ripley", "Ripley", "McNeil", "R Core"),
+   title = c("Exploratory Data Analysis",
+   "Modern Applied Statistics ...",
+   "LISP-STAT", "Spatial Statistics", "Stochastic Simulation",  
+   "Spatial Statistics", "Stochastic Simulation",")
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```r
+ "Interactive Data Analysis", "An Introduction to R"),
+ other.author = c(NA, "Ripley", NA, NA, NA, NA, "Venables & Smith"),
+ stringsAsFactors=FALSE)
> a <- as.scidb(authors)
> b <- as.scidb(books)
> merge(a,b,by.x="surname",by.y="name")[,4:8]

<table>
<thead>
<tr>
<th>nationality</th>
<th>deceased</th>
<th>name</th>
<th>title</th>
<th>other_author</th>
</tr>
</thead>
<tbody>
<tr>
<td>Australia</td>
<td>no</td>
<td>McNeil</td>
<td>Interactive Data Analysis</td>
<td>&lt;NA&gt;</td>
</tr>
<tr>
<td>UK</td>
<td>no</td>
<td>Ripley</td>
<td>Spatial Statistics</td>
<td>&lt;NA&gt;</td>
</tr>
<tr>
<td>UK</td>
<td>no</td>
<td>Ripley</td>
<td>Stochastic Simulation</td>
<td>&lt;NA&gt;</td>
</tr>
<tr>
<td>US</td>
<td>no</td>
<td>Tierney</td>
<td>LISP-STAT</td>
<td>&lt;NA&gt;</td>
</tr>
<tr>
<td>US</td>
<td>yes</td>
<td>Tukey</td>
<td>Exploratory Data Analysis</td>
<td>&lt;NA&gt;</td>
</tr>
<tr>
<td>Australia</td>
<td>no</td>
<td>Venables</td>
<td>Modern Applied Statistics ...</td>
<td>Ripley</td>
</tr>
</tbody>
</table>

> # cf. The standard R data.frame merge function:
> # merge(authors,books,by.x="surname",by.y="name")

The `merge` implementation has some limitations outlined in its man page. Joins on attributes (instead of along coordinate axes) are presently the most limited kinds and outer joins are only supported in certain cases.

### 5.4 Aggregation

The `aggregate` function performs various aggregation operations on SciDB array objects. Aggregation may be defined over array dimensions and/or array attributes very similarly to standard R aggregation syntax, except that aggregation functions must be valid SciDB aggregate expressions, represented as strings.

```r
> # Aggregation example
> df <- as.scidb(iris)
> aggregate(df, by="Species", FUN="avg(Petal_Length), stdev(Petal_Width)")
```

<table>
<thead>
<tr>
<th>Petal_Length_avg</th>
<th>Petal_Width_stdev</th>
<th>Species</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.462</td>
<td>0.1053856</td>
<td>setosa</td>
</tr>
<tr>
<td>4.260</td>
<td>0.1977527</td>
<td>versicolor</td>
</tr>
<tr>
<td>5.552</td>
<td>0.2746501</td>
<td>virginica</td>
</tr>
</tbody>
</table>
```
Although limited support for aggregation by data frame column is available (see the example in this section), the `aggregate` function performs best along SciDB array dimensions. The `aggregate` function also supports single- and multi-dimensional moving window aggregation along array coordinate systems, and 1-d aggregation along consecutive data values in sparse arrays. See the on-line man page for additional details and examples.

### 5.5 Binding new variables to an array

The SciDB package defines the `bind` function to add variables to arrays similar to the R `cbind` function for data frames. However, `bind` can also operate on higher-dimensional arrays. The example below adds a variable named 'prod' to the SciDB array `df` defined in the last example.

```r
> y <- bind(df, "prod", "Petal_Length * Petal_Width")
> head(y, n=3)
```

<table>
<thead>
<tr>
<th>Sepal_Length</th>
<th>Sepal_Width</th>
<th>Petal_Length</th>
<th>Petal_Width</th>
<th>Species</th>
<th>prod</th>
</tr>
</thead>
<tbody>
<tr>
<td>5.1</td>
<td>3.5</td>
<td>1.4</td>
<td>0.2</td>
<td>setosa</td>
<td>0.28</td>
</tr>
<tr>
<td>4.9</td>
<td>3.0</td>
<td>1.4</td>
<td>0.2</td>
<td>setosa</td>
<td>0.28</td>
</tr>
<tr>
<td>4.7</td>
<td>3.2</td>
<td>1.3</td>
<td>0.2</td>
<td>setosa</td>
<td>0.26</td>
</tr>
</tbody>
</table>

### 5.6 Sorting and enumerating factors

Use the `sort` function to sort on a subset of dimensions and/or attribute of a SciDB array object, creating a new sorted array.

Use the `unique` function to return a SciDB array that removes duplicate elements of a single-attribute SciDB input array.

Use the `index_lookup` function along with `unique` to bind a new variable that enumerates unique values of a variable similarly to the R `factor` function.

Examples follow. Note that we use the SciDB `project` function in one example. `project` presents an alternative syntax to the functionally equivalent column subset selection of variables using brackets.

```r
> x <- as.scidb(iris)
> # Sort x by Petal_Width and Species
> a <- sort(x, attributes=c("Petal_Width","Species"))
> head(a, n=3)
```
The *scidb* Package

<table>
<thead>
<tr>
<th>Sepal_Length</th>
<th>Sepal_Width</th>
<th>Petal_Length</th>
<th>Petal_Width</th>
<th>Species</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>4.9</td>
<td>3.1</td>
<td>1.5</td>
<td>0.1</td>
</tr>
<tr>
<td>2</td>
<td>4.9</td>
<td>3.6</td>
<td>1.4</td>
<td>0.1</td>
</tr>
<tr>
<td>3</td>
<td>4.8</td>
<td>3.0</td>
<td>1.4</td>
<td>0.1</td>
</tr>
</tbody>
</table>

> # Find unique values of Species:
> `unique(x$Species)`

```
Species
0 setosa
1 versicolor
2 virginica
```

> # Add a new variable that enumerates factor levels of Species:
> `head(index_lookup(x, unique(x$Species), "Species"))`

<table>
<thead>
<tr>
<th>Sepal_Length</th>
<th>Sepal_Width</th>
<th>Petal_Length</th>
<th>Petal_Width</th>
<th>Species</th>
<th>Species_index</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>5.1</td>
<td>3.5</td>
<td>1.4</td>
<td>0.2</td>
<td>setosa</td>
</tr>
<tr>
<td>2</td>
<td>4.9</td>
<td>3.0</td>
<td>1.4</td>
<td>0.2</td>
<td>setosa</td>
</tr>
<tr>
<td>3</td>
<td>4.7</td>
<td>3.2</td>
<td>1.3</td>
<td>0.2</td>
<td>setosa</td>
</tr>
<tr>
<td>4</td>
<td>4.6</td>
<td>3.1</td>
<td>1.5</td>
<td>0.2</td>
<td>setosa</td>
</tr>
<tr>
<td>5</td>
<td>5.0</td>
<td>3.6</td>
<td>1.4</td>
<td>0.2</td>
<td>setosa</td>
</tr>
<tr>
<td>6</td>
<td>5.4</td>
<td>3.9</td>
<td>1.7</td>
<td>0.4</td>
<td>setosa</td>
</tr>
</tbody>
</table>

### 5.7 Indexing details and missing values

The integer coordinate systems used to index SciDB arrays are similar to R, except that SciDB integer indices may be zero or negative, and are represented by 62-bit signed integers (R indices use either unsigned positive 31-bit integer or positive 52-bit integer-valued double values). This means that SciDB arrays can’t always be indexed by R. In practice, R users should avoid creating SciDB arrays with coordinate systems that extend past R’s limits.

SciDB attribute values within a cell may be explicitly marked missing, indicated by a special SciDB missing code also referred to as a NULL code. SciDB internally supports a large number of possible missing codes. All SciDB missing code values are mapped to `NA` values in R.
6 Lazy evaluation, array promises, and garbage collection

Most operations on SciDB array objects return array promises—new SciDB array objects that have not been fully evaluated by SciDB yet, but that promise to return values when asked. SciDB array promises are essentially just SciDB query expressions together with a schema that the resulting output array will have once it’s been evaluated and an environment providing context for the evaluation. In such cases, the name slot of a scidb or scidbdf object shows the SciDB query expression.

Sometimes it can be more efficient to explicitly evaluate and cache a SciDB array then to use it as an array promise. This can always be achieved by using the scidbeval function, or by using the `eval=TRUE` argument in the functions that support that. Dynamically allocated arrays use a naming convention that begins with “R_array” and end with a unique numeric identifier determined by the current SciDB session.

Ephemeral intermediate arrays are by default connected to R’s garbage collector and automatically deleted from the SciDB catalog when they are no longer needed by R. Users can disconnect SciDB array objects from R’s garbage collector (making stored arrays persistent in SciDB), by using the gc=FALSE flag in the as.scidb and scidbeval functions.

SciDB arrays keep references to other SciB array dependencies in a list within an environment, preventing them from automatic garbage collection so that the promise can be evaluated. The environment is slot is named @gc in the array objects and the dependency list is @gc$depend.

Here is an example:

```r
> # Create a 5x4 SciDB matrix named 'test' in the database. We set `gc=TRUE` to make it ephemeral
> A <- as.scidb(matrix(rnorm(20), nrow=5), name="test", gc=TRUE)
> A@name
[1] "test"

> # Assign the third row of the transpose to a scidb object B:
> B <- t(A)[3,]
> # B is a promise object--it's name is an (unevaluated) SciDB query expression
> # that depends on A. But it has a valid output schema, and we also see that
> # it depends ultimately on the SciDB array 'test':
> B@name
```
The scidb Package

[1] "redimension(redimension(between(transpose(test), 3, null, 3, null), <val:double NULL>  
> B@schema
[1] "<val:double NULL DEFAULT null> [i=0:4,5,0]"
> B@gc$depend[[1]]@gc$depend[[1]]@name
[1] "transpose(test)"
> # We can use `scidbeval` to force B's evaluation and storage inside SciDB:
> C <- scidbeval(B)
> # C has the same schema as B, but is now evaluated and stored inside SciDB
> # using an automatically-generated name. Unlike B, C does not depend on
> # the `test` array any more.
> C@name

[1] "R_array2bb38c448db1101074813220"
> C@schema
[1] "<val:double NULL DEFAULT null> [i=0:4,5,0]"

7 SciDB-specific functions

This section briefly summarizes a number of utility functions that are simple wrappers of miscellaneous SciDB operators. Except for count, which returns a number, the listed functions return references to new SciDB array objects.

- attribute_rename Rename one or more attributes of a SciDB array.
- build Creates new dense arrays from SciDB expressions and schema.
- cast Create a new array by altering the schema of an array.
- count Return the number of nonempty cells of an array.
- cumulate Compute a cumulative aggregate function, for example a cumulative sum or product.
- dimension_rename Rename one or more dimensions of a SciDB array.
The `scidb` Package

- **index_lookup** Look up attribute values in a mapping array, adding a new attribute with the index.
- **project** Pick out one or more attributes from an array.
- **redimension** Redimension an array into the provided new schema.
- **regrid** Decimate an array by binning it along one or more dimensions and applying an aggregate function to the bins.
- **rename** Rename a SciDB array.
- **repart** Repartition an array.
- **reshape** Reshape an array.
- **schema** Show the schema of a SciDB array object.
- **scidblist** List SciDB arrays (also `scidbls`).
- **scidbremove** Delete SciDB arrays (also `scidbrm`).
- **slice** Slice a SciDB array along one of its axes, returning a lower-dimensional array.
- **sort** Sort the elements of an array.
- **subarray** Perform a SciDB subarray or between operation on an array.
- **unique** Return a set of unique elements of a sorted array.
- **unpack** Unpack a SciDB array into a 1-d table.
- **xgrid** Prolong one or more coordinate axes by replicating cell values.

See the online manual pages for more details and examples.

8 Miscellaneous functions

This catch-all section summarizes miscellaneous functions.

8.1 Histograms

The generic function `hist` computes a histogram of the given data values in a SciDb array object. If `plot=TRUE`, the resulting object of class histogram is plotted by `plot.histogram`, before it is returned.

This histogram function only supports equidistant breaks. If `right=TRUE` (default), the histogram cells are intervals of the form ‘(a, b]’, i.e., they include their right-hand endpoint, but not their left one.
8.2 Last-value imputation

The `na.locf` function is a generic function for replacing each missing or empty value with the most recent non-missing value prior to it.

Unlike the usual `na.locf` function from the zoo package, the SciDB `na.locf` function fills in both missing (SciDB null values) and empty (SciDB sparse) values with the last non-missing and non-sparse value along the indicated dimension.

Time series represented in SciDB are often sparse arrays. The `na.locf` function defines a convenient way to fill in all missing values along the time coordinate axis down to the time resolution.

Caution! The output array is a mostly dense, filled-in version of the input array. If the time resolution is very fine and the input array very sparse, then the output array can be huge. Consider using regrid first on very fine time scales to reduce their resolution.

The default `na.locf` method in the zoo package unfortunately overrides this function (it uses ANY in its method signature). If you need to use SciDB arrays and the zoo package, prefix SciDB’s version with `scidb::na.locf(...)`.

9 Running SciDB queries

The `iquery` function executes SciDB queries using either the SciDB array functional language (AFL) or declarative array query language (AQL) syntax. When AFL is used, the `iquery` function optionally returns query results in an R data frame if the argument `return=TRUE` is specified. Returned output is similar to output obtained by the SciDB `iquery` command-line program with the `-olcsv+` option. The `iquery` function does not return anything by default.

Query results returned by the `iquery` function are internally presented to R using a generic CSV format, providing very flexible support for many data types. (The $n$-dimensional array class described in the next section uses a binary data exchange method between R and SciDB.) Note that, although R and SciDB have a number of common data types, each system contains types not supported by the other. Thus, conversion errors may arise. The `iquery` function is designed to reasonably minimize such issues and simplify basic data transfer between the systems. Data types common to R and SciDB include double-precision numeric, character string,
logical, and 32-bit integers. The iquery function supports standard R read.table parameter options to facilitate type conversion.

The following code example illustrates basic use of iquery.

```r
> # Manually build a 1-d SciDB array named "P:"
> iquery("store(build(<x:double>[i=0:99,100,0],asin(1)*i/25),P)"
> # Return a query as a data.frame, manually specifying returned column classes:
> S <- iquery("apply(P,y,sin(x))",return=TRUE,
+   colClasses=c("integer","double","complex"))
> head(S, n=3)

   i   x             y
1 0 0.00000000 0.00000000+0i
2 1 0.06283185 0.06279052+0i
3 2 0.12566371 0.12533323+0i

9.1 Iterating over query results

The iquery function returns query results into a single R data frame by default. Large results expected to contain lots of rows may be iterated over by setting the iterative=TRUE argument. When iquery is used with the iterative=TRUE setting, it returns an iterator that iterates over chunks of rows of the result data frame. Iterators are defined by the iterators package. Their data may be directly accessed with the nextElem method, or indirectly with foreach. See the iterators and foreach packages for many examples and further documentation of their use.

```r
> # Build a small 1-d SciDB test array:
> iquery("store(build(<x:double>[i=1:10,10,0],i/10.0),A)"
> # Return the result of a SciDB apply operator in an R iterator with a
# block size of at most 7 rows at a time:
> library("iterators")
> it <- iquery("apply(A,y,sqrt(x))", return=TRUE, iterative=TRUE, n=7)
> nextElem(it)

   i   x     y
1 1 0.1 0.3162278
```
$ \%<\% \quad \text{abs} \quad \text{acos} \quad \text{aggregate} \\
[ \quad \text{antijoin} \quad \text{apply} \quad \text{as.scidb} \\
\text{all.equal} \quad \text{atan} \quad \text{attribute.rename} \quad \text{between} \\
\text{asin} \quad \text{build} \quad \text{c} \quad \text{cast} \\
\text{bind} \quad \text{colnames} \quad \text{colnames<-} \quad \text{cos} \\
\text{cbind} \quad \text{crossprod} \quad \text{cumulate} \quad \text{default} \\
\text{count} \quad \text{diag} \quad \text{diff} \quad \text{dim}$
12 Package installation

Installation proceeds in two steps: installing the R package on any computer that has a network connection to a SciDB database, and installing a simple network service on the SciDB database coordinator computer.

12.1 Installing the R package from sources on GitHub

The scidb package source is maintained in the SciDBR GitHub repository. That’s where the most up-to-date version of the package is available. Released versions of the package posted to CRAN are updated much less frequently, approximately semiannually. A git tag indicates each CRAN release version of the package in the source code repository.

The wonderful devtools R package makes installation of source packages from GitHub nearly as simple as installation from CRAN.
library("devtools")
install_github("SciDBR", "Paradigm4", quick=TRUE)

12.2 Installing the R package from CRAN

The scidb package is available on CRAN. Start an R session and run:

install.packages("scidb")

12.3 Installing the simple HTTP service for SciDB

The SciDB R package requires installation of a simple open-source HTTP network service called shim on the computer that SciDB is installed on. The service needs to be installed only on the SciDB coordinator computer, not on client computers that connect to SciDB from R. It’s available in packaged binary form for supported SciDB operating systems, and as source code which can be compiled and deployed on any SciDB installation.

Both installation approaches install the shim network service on the SciDB coordinator computer. Installing as a service requires root permission. The compiled source code version requires no special permissions to run.

Installation from binary software packages for SciDB-supported operating systems is easiest. Detailed up-to-date information can be found on Paradigm4’s laboratory on Paradigm4’s Github repository at https://github.com/Paradigm4/shim/wiki/Installing-shim. We outline installation for each supported operating system below. See our github page for source code. The open source package author, Bryan Lewis, maintains binary packages for SciDB-supported operating systems. They are tied to specific versions of SciDB. The present version is 13.12 (December, 2013).

12.3.1 Installation on RHEL/CentOS 6

# Install with:
wget http://paradigm4.github.io/shim/shim-13.12-1.x86_64.rpm
rpm -i shim-13.12-1.x86_64.rpm

# (Uninstall, if desired, with:)
yum remove shim
12.3.2 Installation on Ubuntu 12.04

# Install with:
wget http://paradigm4.github.io/shim/shim_13.12_amd64.deb
sudo gdebi shim_13.12_amd64.deb

# (Uninstall, if desired, with:)
apt-get remove shim

See the Wiki and web pages at https://github.com/Paradigm4/shim/ for up to date package information and source code.

The installed shim network service exposes SciDB as a very simple HTTP API. It includes a simple browser-based status and query tool. After installing shim, point your browser to the I.P. address of the SciDB coordinator machine and port 8080. Note that this API is not official and may change in the future. Help drive those changes by contributing ideas, code and bugfixes to the project on Github, or feel free to discuss the service on the SciDB.org/forum or via Github issues.

12.4 Package options, miscellaneous notes, and software license

The scidb package defines several global package options. Package options may be set and retrieved with the R options function, and are listed in Table 2.

<table>
<thead>
<tr>
<th>Option</th>
<th>Default value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>scidb.debug</td>
<td>NULL</td>
<td>Set to TRUE to display all queries issued to the SciDB engine and other debugging information.</td>
</tr>
<tr>
<td>scidb.index.sequence.limit</td>
<td>100 000 000</td>
<td>Maximum allowed scidb array object sequential indexing limit (for larger ranges, use between)</td>
</tr>
<tr>
<td>scidb.max.array.elements</td>
<td>100 000 000</td>
<td>Maximum allowed non-empty elements to return in a subsetting operation of a scidb array object.</td>
</tr>
</tbody>
</table>

Package options
Miscellaneous notes:

- R does not support 64-bit integer types. 64-bit signed and unsigned integers smaller than $2^{53}$ in magnitude will be represented as double-precision floating point numbers. 64-bit integers outside that range appear as $+/-\infty$. All other integers (int8, uint8, int16, uint16, etc.) are represented in R by 32-bit signed integers. The SciDB uint32 type is not supported.

- R doesn’t support single-precision floating point numbers. iquery results convert single-precision numbers within SciDB to double-precision floating-point numbers in R. Single-precision SciDB numbers are not supported by the scidb array class.

- SciDB does not natively support complex numbers. Loading complex numbers directly into SciDB from R is not defined.

- The iquery function provides the most flexible mechanism for type conversion between the systems, fully under user control using read.table options.

- Naming convention and associated restrictions vary between R and SciDB. For example, SciDB does not allow decimal points in attribute names. The package may alter names with character substitution to reconcile names when it is reasonable to do so. A warning is emitted whenever an object is automatically renamed in this way.

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