Multiple-table data in R

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Abstract

Data frames are integral to R. They provide a standard format for passing data to model-fitting and plotting functions. This standard makes it easier for experienced useRs to learn new functions, because most developments in R continue to accept a single data frame as input. Still, many data sets do not easily fit into a single data frame, and my field of community ecology provides many examples of such inherently multiple-table data (e.g. fourth-corner problem and other trait-based data sets). Storing such data in a single data frame results in either large numbers of meaningless missing values or storage of redundant information. These storage problems have led ecologists to model summaries of their data (e.g. community-weighted trait matrices), rather then their data itself. Perhaps more importantly, my experience with manipulating such data using data frames has resulted in difficult-to-read workflows with many lines. The multitable package introduces new data storage objects called data.lists, which are extensions of data.frames. As data.lists can be coerced to data.frames, they can be used with all R functions that accept an object that is coercible to a data.frame (e.g. lm; plot; lme; and many more). The multitable package also provides several mechanisms for simplifying the manipulation of data.list objects.

1 Introduction

The standard data management paradigm in R is based on data.frame objects, which are two-dimensional data tables with rows and columns representing replicates and variables respectively. Standard R workflows require that all of the data to be analyzed are organized into a single data frame, and hypotheses about the relationships between variables in the data frame are expressed using formula objects; data frames and formulas are combined by passing them to functions that produce analyses (e.g. plots; fitted models; summary statistics) [1]. This framework allows scientists to concentrate on their primary interests—the relationships between variables—without explicit reference to complex mathematical and algorithmic details. It also provides access to those details, which are required (1) for more effective analyses and (2) to develop new methods of analysis within the framework. As new methods are developed, researchers simply pass their data frames to new functions in much the same way they

would pass them to older functions. Thus, by separating low-level methods development from high-level data analysis, R fosters the formation of a community of researchers where both methodologists and analysts can have mutually beneficial interactions.

However, research in my field of community ecology has led my colleagues and I to data sets that do not easily fit within a single data frame. A common example is the fourth-corner problem [5], in which three data tables are to be analyzed: a sites-by-species table of abundances or occurrences; a table of environmental variables at each site; and a table of traits for each species (Fig. 1). Such data are characterized by a conspicuous (lower-right) 'fourth-corner', where there are no data. These fourth-corners of missing data are not caused by the usual problems (e.g. broken field equipment; budget restrictions; bad weather; dead subjects), but are part of the study design itself. The fourth-corner problem is a special case of a general 'multiple-table problem', which can be much more complex (e.g. could involve three-dimensional 'cubes' of data, Fig. 2). The challenge of analyzing such multiple-table data sets in R is that it is not obvious how to organize them into a single data.frame, which is required in standard R workflows. Our goal with the multitable package is to provide tools for analyzing multiple-table data sets within this standard R framework.

One possible solution is to develop new R analysis functions—or new software packages altogether—that are specifically designed to accept several tables as input. There has been a fair amount of work in this direction, focusing on data with a fourth-corner problem [2, 5, 3, 7, 6, 4]. However, this work does not apply to data sets that have other more complex multiple-table data structures (e.g. zooplankton communities in Lac Croche, Fig. 2; Ref??). One approach to such issues would be to build new data analysis functions for each new data structure. But such an approach is less than ideal, as it would require that new methods be learned for each new structure—it does not take advantage of the large number of tools developed within the standard R framework of data frames and formulas. The multitable package provides an alternative approach, by introducing a multiple-table generalization of data frames—called data lists—which can be analyzed with virtually any function that can be used to analyze a data frame. Thus, instead of providing new methods of analysis, multitable provides new methods of data management.

There are several existing R packages that are designed to make data management easier (e.g. reshape2; (author?), (year?)). In fact, the mefa and mefa4 packages have been developed to organize data with a slight generalization¹ of the fourth-corner problem [8]. The multitable package has much in common with mefa, but there are noticeable differences. For example, mefa provides more extensive tools for data summarization than multitable and mefa4 integrates tools for sparse-matrix computations. On the other hand, multitable is designed to handle more general data structures than mefa or mefa4 (e.g. mefa cannot organize the Lac Croche data structure, Fig. 2). However,

¹Several community matrices—called segments—with identical dimensions are allowed in mefa.

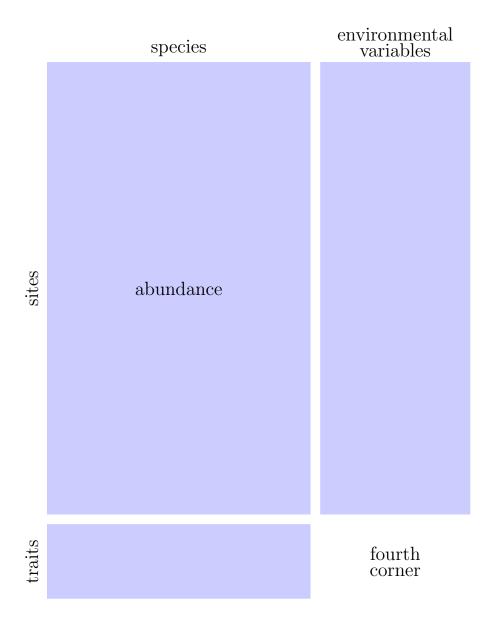
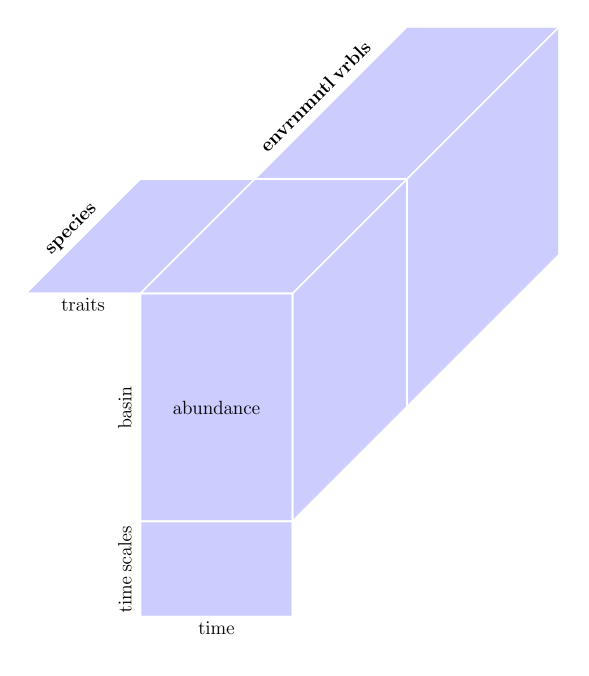


Figure 1: Fourth corner problem.



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Figure 2: The structure of the Lac Croche zooplankton community data.

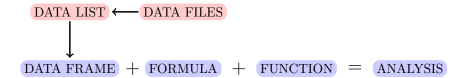


Figure 3: The multitable paradigm for including multiple-table data (in red) into the standard R workflow (in blue). Data lists are used to organize and manipulate multiple-table data as a single R object, even though it must be stored in multiple text-based data files. When such data are required for analysis, they are coerced into a data frame. Once in data frame form, they can be used in analyses by combining them with formulas (to specify hypothetical relationships between variables) and functions (to call computational methods).

we hope that mefa and multitable will be complementary, not competitive.

The multitable model of data organization is illustrated in Figure 3 (mefa uses a similar model). The elements of the standard R workflow are in blue: data frames; formulas; functions; and analyses. The multitable package seeks to facilitate the use of such workflows with multiple-table data by creating tools (arrows and red boxes) for organizing and manipulating such data. These tools are based on a new kind of object, called a data list, which is used to organize multiple-table data. Data lists can be manipulated much like data frames (e.g. variables can be transformed; groups of observations extracted or removed). Our design principle was to keep the manipulation of data lists as similar as possible to the manipulation of data frames. Once data lists are ready for analysis multitable provides tools for coercing them into data frames, thereby entering the standard R workflow. Importantly, data, formulas, and functions are kept separate, thus preserving the benefits of using R in this standard way.

The purpose of this vignette is to justify and introduce the use of the multitable package. I begin by describing the structure of an example data.list object. Then I illustrate one of the most powerful features of data.lists: methods that allow related variables, which cannot fit into a single data frame, to be subscripted simultaneously. Next I show that variables in data lists can be transformed and modeled, in the same manner that is standard for variables in data frames. Finally, I describe a simple method for creating a data list of your own data, and use this method to introduce some useful concepts associated with multiple-table data.

2 The structure of data lists

The multitable package comes with a fictitious data.list, to illustrate how these objects work.

```
> library(multitable)
```

- > data(fake.community)
- > fake.community

abundance:

, , capybara

	2009	2008	1537
${\tt midlatitude}$	4	0	0
subtropical	0	10	0
tropical	8	0	0
equatorial	0	7	0
arctic	0	0	0
subarctic	0	0	0

, , moss

	2009	2008	1537
midlatitude	0	6	0
subtropical	0	0	0
tropical	9	0	0
equatorial	0	3	0
arctic	5	0	0
subarctic	0	0	0

, , vampire

	2009	2008	1537
midlatitude	0	0	0
subtropical	0	0	1
tropical	0	0	0
equatorial	0	0	0
arctic	0	0	0
subarctic	0	0	0

Replicated along: || sites || years || species ||

${\tt temperature:}$

2009 2008 1537

```
midlatitude NA 10
                     NA
subtropical 25 20
                     NA
          48 50
tropical
                     NA
equatorial
            50 30
                     NA
arctic
           -37 -30
                     NA
subarctic
            3
                0
                     NA
Replicated along: || sites || years ||
precipitation:
          2009 2008 1537
midlatitude NA 20
                     NA
subtropical 99 100
                     NA
tropical 149 150
                     NA
equatorial 199 200
                     NA
         21 20
arctic
                     NA
subarctic
          41 40
                     NA
Replicated along: || sites || years ||
body.size:
capybara
          moss vampire
    140
           NA
                    190
Replicated along: || species ||
metabolic.rate:
        moss vampire
capybara
     20
           5
Replicated along: || species ||
homeotherm:
capybara
           moss vampire
     Y
           N
Levels: N Y
Replicated along: || species ||
REPLICATION DIMENSIONS:
 sites years species
     6
           3
```

At first sight, this data.list object looks very different from standard data.frame objects, but on second look we can see that they are really quite similar. Just like data frames, data lists are composed of a number of variables—in this case, we have six variables (abundance; temperature; precipitation; body size; metabolic rate; and homeotherm) each identified in the printed object above by underlined names. The variables in data lists must be printed in this sequential manner, rather than as columns neatly lined up in a data frame, precisely because the variables in multiple-table data sets do not line up neatly; this is the problem multitable seeks to address.

Also as in data frames, the replication of variables in data lists are represented as vectors of values. The main difference between the two objects in this regard is that the vectors that represent variables in data lists have a dim (i.e. dimension) attribute, which gives it further structure. In R, vectors with dim attributes are best thought of as matrices and arrays of numbers. For example, the abundance variable is replicated along three dimensions (sites; years; and species), and therefore is a three dimensional array of data. This information is indicated above after the variable itself is printed. Some variables are only replicated along two dimensions (e.g. temperature and precipitation) and others only a single dimension (e.g. body size; metabolic rate; and homeotherm).

Importantly however, although the variables are not replicated along all of the same dimensions, they do share dimensions; and it is this dimension sharing that allows us to relate variables to each other. To appreciate the dimension sharing of this example, we can use the summary method for data.list objects.

> summary(fake.community)

	abundance	temper	ature	precipi	tation	body.size
sites	TRUE		TRUE		TRUE	FALSE
years	TRUE		TRUE		TRUE	FALSE
species	TRUE		FALSE		FALSE	TRUE
	${\tt metabolic}.$	rate h	omeoth	nerm		
sites	F	ALSE	FI	ALSE		
years	F	ALSE	FI	ALSE		
species		TRUE	7	ΓRUE		

This method returns a logical table with dimensions of replication as rows and variables as columns. A value of TRUE appears in cells corresponding to variables that are replicated along a particular dimension, and a value of FALSE appears otherwise. We can see that the sites and years dimensions relate abundance, temperature, and precipitation; whereas, the species dimension relates abundance, body size, metabolic rate, and homeotherm.

Note that some FALSE entries are bio-physical necessities, whereas some are properties of the study design. For example, suppose that later in the study, the researchers decided that it was necessary to get some idea of the spatial variation in metabolic rates. It would then be possible to measure metabolic rates of the species at different sites, thereby changing the FALSE associated

with the metabolic rate-sites cell to a TRUE. To the contrary, it is physically and logically impossible to measure the precipitation of a species, and so this FALSE is necessarily FALSE.

3 Subscripting data lists

This structure relating variables and dimensions of replication, allows us to manipulate multiple variables simultaneously. In particular, multitable makes it possible to extract pieces of a data list while maintaining its structure. For example, examining the data suggests that 1537 might have been an outlying year relative to 2008 and 2009. We can exclude data from 1537 just as we would with a single R array.

```
> fake.community[,c("2008","2009"),]
abundance:
_____
, , capybara
            2008 2009
midlatitude
               0
subtropical
              10
                    0
tropical
equatorial
               7
                    0
arctic
               0
                    0
subarctic
                    0
, , moss
            2008 2009
midlatitude
               6
                    0
subtropical
               0
               0
                    9
tropical
               3
                    0
equatorial
                    5
arctic
               0
subarctic
               0
                    0
, , vampire
            2008 2009
midlatitude
               0
                    0
subtropical
               0
tropical
               0
                    0
                    0
equatorial
               0
               0
                    0
arctic
subarctic
```

```
Replicated along: || sites || years || species ||
temperature:
          2008 2009
midlatitude 10 NA
subtropical 20 25
          50 48
tropical
equatorial 30 50
arctic
           -30 -37
          0
subarctic
               3
Replicated along: || sites || years ||
precipitation:
         2008 2009
midlatitude 20 NA
subtropical 100 99
tropical 150 149
equatorial
           200 199
      20 21
ic 40 41
arctic
subarctic
Replicated along: || sites || years ||
body.size:
-----
capybara moss vampire
   140
          NA 190
Replicated along: || species ||
metabolic.rate:
capybara
         moss vampire
          5
Replicated along: || species ||
homeotherm:
-----
capybara moss vampire
      Y
           N
```

```
Levels: N Y
Replicated along: || species ||
REPLICATION DIMENSIONS:
sites years species
6 2 3
```

This command returns the same data list of variables but without the data from 1537. Note that every variable replicated along the years dimension is subscripted appropriately, while variables that are not replicated along this dimension are unchanged. As another example, perhaps we want all of the data on the first species (i.e. capybara) in 1537 for the first three sites.

```
> fake.community[1:3,"1537",1]
abundance:
, , capybara
            1537
midlatitude
             0
subtropical
              0
tropical
               0
Replicated along: || sites || years || species ||
temperature:
           1537
midlatitude
             NA
subtropical
             NA
tropical
             NA
Replicated along: || sites || years ||
precipitation:
-----
           1537
midlatitude
             NA
subtropical
             NA
tropical
             NA
Replicated along: || sites || years ||
body.size:
```

```
capybara
     140
Replicated along: || species ||
metabolic.rate:
capybara
      20
Replicated along: || species ||
homeotherm:
-----
capybara
       Y
Levels: N Y
Replicated along: || species ||
REPLICATION DIMENSIONS:
  sites
         years species
```

Notice also that for each different subset of the data, the new replication dimensions are printed after the data.

4 Transforming variables in data lists

Often we need to transform variables before passing data frames to functions. This is easily done with variables in data lists as well. For example, suppose we want to make a $\log(x+1)$ transformation of the abundance data.

, , moss

```
2009
                          2008 1537
midlatitude 0.000000 1.945910
                                  0
subtropical 0.000000 0.000000
                                  0
            2.302585 0.000000
                                  0
tropical
            0.000000 1.386294
                                  0
equatorial
            1.791759 0.000000
                                  0
arctic
subarctic
            0.000000 0.000000
                                  0
```

, , vampire

```
2009 2008
                            1537
                    0 0.0000000
midlatitude
               0
subtropical
               0
                    0 0.6931472
tropical
               0
                    0 0.0000000
equatorial
               0
                    0 0.0000000
arctic
               0
                    0 0.000000
               0
                    0 0.0000000
subarctic
```

```
attr(,"subsetdim")
sites years species
TRUE TRUE TRUE
```

We note that fake.community has a lot of missing values, which were useful for illustrating how data lists handle missing values, but will make further illustrations somewhat underwhelming. We can replace these missing values with values using the standard logic of R replacement.

5 Simple analysis functions

Data lists can be passed 'as is' to many standard functions in R that normally take data frames. In the next section I will define this class of functions in more detail, but for now consider this simple example. Perhaps we want to explore whether the interaction between body size and temperature has an influence on abundance. As a first attempt at model building, we fit a linear model using lm.

```
> lm(abundance ~ (body.size*temperature),data=fake.community)
```

```
Call:
    lm(formula = abundance ~ (body.size * temperature), data = fake.community)
    Coefficients:
              (Intercept)
                                        body.size
                4.484e-01
                                       -1.718e-03
              temperature body.size:temperature
                3.634e-03
                                        5.041e-07
And this works just as well with mixtures of categorical and numerical data.
    > lm(abundance ~ -1+(homeotherm*temperature),data=fake.community)
    lm(formula = abundance ~ -1 + (homeotherm * temperature), data = fake.community)
    Coefficients:
                homeothermN
                                          homeothermY
                   0.228770
                                              0.318948
                temperature homeothermY:temperature
                   0.001186
                                              0.007512
It also works with other 'simple' functions, such as rlm (robust linear model) in
the MASS package.
    > library(MASS)
    > rlm(abundance ~ (body.size*temperature),data=fake.community)
    rlm(formula = abundance ~ (body.size * temperature), data = fake.community)
    Converged in 10 iterations
    Coefficients:
              (Intercept)
                                       body.size
             2.606699e-05
                                   -1.076827e-07
              temperature body.size:temperature
             3.043212e-07
                                   -8.994997e-10
    Degrees of freedom: 51 total; 47 residual
      (3 observations deleted due to missingness)
    Scale estimate: 5.26e-05
```

Therefore, in many cases, data lists enter the standard R workflow in exactly the same manner as data frames.

6 Coercing data lists to data frames

The reason that unmodified data lists can be passed to some functions that are expecting data frames, is that these functions try to coerce whatever data object

they receive into a data frame. When the multitable package is loaded, these functions can find a method for making such a conversion. This method can be accessed by users directly via the as.data.frame function from the R base package. For example, we can pass the fake.community data to as.data.frame.

- > fake.community.df <- as.data.frame(fake.community)</pre>
- > fake.community.df

- Tano. Community . al				
		=	${\tt precipitation}$	=
midlatitude.2009.capybara		NA	NA	140
subtropical.2009.capybara	0.0000000	25	99	140
tropical.2009.capybara	2.1972246	48	149	140
equatorial.2009.capybara	0.0000000	50	199	140
arctic.2009.capybara	0.0000000	-37	21	140
subarctic.2009.capybara	0.0000000	3	41	140
midlatitude.2008.capybara		10	20	140
subtropical.2008.capybara	2.3978953	20	100	140
tropical.2008.capybara	0.0000000	50	150	140
equatorial.2008.capybara	2.0794415	30	200	140
arctic.2008.capybara	0.0000000	-30	20	140
subarctic.2008.capybara	0.0000000	0	40	140
midlatitude.1537.capybara		5	5	140
subtropical.1537.capybara	0.0000000	10	50	140
tropical.1537.capybara	0.0000000	30	75	140
equatorial.1537.capybara	0.0000000	20	50	140
arctic.1537.capybara	0.0000000	-80	2	140
subarctic.1537.capybara	0.0000000	-10	7	140
midlatitude.2009.moss	0.0000000	NA	NA	1
subtropical.2009.moss	0.0000000	25	99	1
tropical.2009.moss	2.3025851	48	149	1
equatorial.2009.moss	0.0000000	50	199	1
arctic.2009.moss	1.7917595	-37	21	1
subarctic.2009.moss	0.0000000	3	41	1
midlatitude.2008.moss	1.9459101	10	20	1
subtropical.2008.moss	0.0000000	20	100	1
tropical.2008.moss	0.0000000	50	150	1
equatorial.2008.moss	1.3862944	30	200	1
arctic.2008.moss	0.0000000	-30	20	1
subarctic.2008.moss	0.0000000	0	40	1
midlatitude.1537.moss	0.0000000	5	5	1
subtropical.1537.moss	0.0000000	10	50	1
tropical.1537.moss	0.0000000	30	75	1
equatorial.1537.moss	0.0000000	20	50	1
arctic.1537.moss	0.0000000	-80	2	1
subarctic.1537.moss	0.0000000	-10	7	1
midlatitude.2009.vampire	0.0000000	NA	NA	190

subtropical.2009.vampire	0.0000000	25	99
tropical.2009.vampire	0.0000000	48	149
equatorial.2009.vampire	0.0000000	50	199
arctic.2009.vampire	0.0000000	-37	21
subarctic.2009.vampire	0.0000000	3	41
midlatitude.2008.vampire	0.0000000	10	20
subtropical.2008.vampire	0.0000000	20	100
tropical.2008.vampire	0.0000000	50	150
equatorial.2008.vampire	0.0000000	30	200
arctic.2008.vampire	0.0000000	-30	20
subarctic.2008.vampire	0.0000000	0	40
midlatitude.1537.vampire	0.0000000	5	5
subtropical.1537.vampire	0.6931472	10	50
tropical.1537.vampire	0.0000000	30	75
equatorial.1537.vampire	0.0000000	20	50
arctic.1537.vampire	0.0000000	-80	2
subarctic.1537.vampire	0.0000000	-10	7
Subarcule.1007.vampire	metabolic.rate		,
midlatitude.2009.capybara	20		
subtropical.2009.capybara	20		
tropical.2009.capybara	20		
	20		
equatorial.2009.capybara			
arctic.2009.capybara	20		
subarctic.2009.capybara	20		
midlatitude.2008.capybara	20		
subtropical.2008.capybara	20		
tropical.2008.capybara	20		
equatorial.2008.capybara	20		
arctic.2008.capybara	20		
subarctic.2008.capybara	20		
midlatitude.1537.capybara	20		
subtropical.1537.capybara	20		
tropical.1537.capybara	20		
equatorial.1537.capybara	20		
arctic.1537.capybara	20		
subarctic.1537.capybara	20		
midlatitude.2009.moss	5		
subtropical.2009.moss	5	N	
tropical.2009.moss	5		
equatorial.2009.moss	5		
arctic.2009.moss	5		
subarctic.2009.moss	5		
midlatitude.2008.moss	5		
subtropical.2008.moss	5		
tropical.2008.moss	5		
equatorial.2008.moss	5	N	

arctic.2008.moss	5	N
subarctic.2008.moss	5	N
midlatitude.1537.moss	5	N
subtropical.1537.moss	5	N
tropical.1537.moss	5	N
equatorial.1537.moss	5	N
arctic.1537.moss	5	N
subarctic.1537.moss	5	N
midlatitude.2009.vampire	0	N
subtropical.2009.vampire	0	N
tropical.2009.vampire	0	N
equatorial.2009.vampire	0	N
arctic.2009.vampire	0	N
subarctic.2009.vampire	0	N
midlatitude.2008.vampire	0	N
subtropical.2008.vampire	0	N
tropical.2008.vampire	0	N
equatorial.2008.vampire	0	N
arctic.2008.vampire	0	N
subarctic.2008.vampire	0	N
midlatitude.1537.vampire	0	N
subtropical.1537.vampire	0	N
tropical.1537.vampire	0	N
equatorial.1537.vampire	0	N
arctic.1537.vampire	0	N
subarctic.1537.vampire	0	N

The resulting data frame contains one column for each variable and one row for each combination of replicates across the three dimensions of replication. Notice that the row names are automatically generated to be informative about the dimensions of replication that have been collapsed into a single dimension. Unlike the corresponding data list object, the data frame has redundancy. For example, because the traits are only replicated along species there are only three unique trait values, one for each of the three species. These three values are repeated so that all of the variables can be stored side-by-side in a single data frame.

By storing these data in a single data frame, we can now pass them to any function that accepts data frames. For example, we can graphically examine the interaction between an environmental variable and a trait using the xyplot function from the lattice package (Fig. 4). Because these are completely fake data I won't make too much out of the results, but there doesn't seem to be much of an interaction between body size and temperature. The exciting thing about this graph is that it suggests a general graphical method for exploring the interactions between traits and environmental variables on community composition.

```
> library(lattice)
> xyplot(abundance ~ temperature | body.size,data=fake.community.df)
```

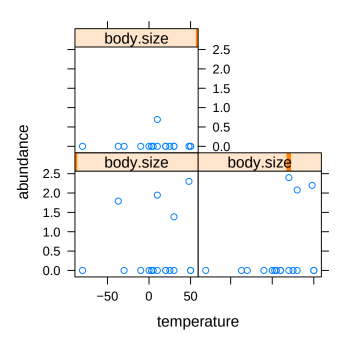


Figure 4: xyplot

7 How data lists are made

Up until now we have used an existing data list to illustrate the use of the multitable package. Although there are several ways to create data lists, there is one way that provides the simplest framework for understanding the difference between variables and dimensions of replication, which is an important distinction to understand in order to use multitable most effectively.

Consider a data frame of species abundances counted at various sites.

> abundance

	sites	species	${\tt abundance}$
1	${\tt midlatitude}$	capybara	4
2	subtropical	capybara	10
3	tropical	capybara	8
4	equatorial	capybara	7
5	arctic	moss	5
6	${\tt midlatitude}$	moss	6
7	tropical	moss	9
8	equatorial	moss	3
9	subtropical	vampire	1

We have six sites and three species, but each species is not present at each site and so there are missing site-species combinations. Related to this data frame we have a data frame of environmental variables at each site and a data frame of traits for each species.

> environment

	site	es tempera	ature	precipitat	tion
1	subarcti	ic	0		40
2	midlatitud	le	10		20
3	subtropica	al	20		100
4	tropica	al	50		150
5	equatoria	al	30		200
>	trait				
	species b	oody.size	metal	oolic.rate	
1	capybara	140		20	
2	moss	5		5	
3	vampire	190		0	

To make things interesting to scientists with real data, we assume that our environmental data are missing from the arctic site (perhaps because it is too harsh and remote).

The three data frames are related because they share two columns: sites and species. The specific pattern of sharing for these data can be illustrated with a bipartite graph (i.e. matching diagram; Fig. 5). Columns that are shared between data frames are called *dimensions of replication* and those that are not are called *variables*. The reason for this terminology is that in standard

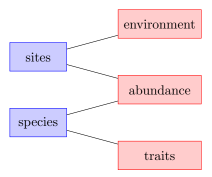


Figure 5: Bipartite graph of the multiple-table structure of data with a standard fourth-corner problem (Fig. 1). Dimensions of replication are in blue and tables are in red.

single-table statistical settings, we are able to relate variables because they are replicated along some common dimension. For example, we could relate pH and temperature if they were both replicated along the same set of lakes. Similarly, we can relate the variables in several tables together if they share columns (i.e. dimensions of replication).

To create a data list out of these data frames we use the dcast function, which was inspired by the acast function in the reshape2 package [9].

```
> dl <- dlcast(list(abundance,environment,trait),</pre>
         dimids=c("sites", "species"),
         fill=c(0,NA,NA)
)
> d1
abundance:
            capybara moss vampire
midlatitude
                    4
                         6
                                  0
                   10
subtropical
                         0
                                  1
                                  0
tropical
                    8
                         9
                    7
equatorial
                         3
                                  0
                    0
                         5
arctic
                                  0
subarctic
                    0
                         0
                                  0
Replicated along: || sites || species ||
temperature:
  -----
midlatitude subtropical
                            tropical
                                       equatorial
                                                        arctic
                                   50
                                                30
         10
                                                            NA
  subarctic
          0
```

```
Replicated along: || sites ||
precipitation:
midlatitude subtropical
                            tropical
                                       equatorial
                                                        arctic
         20
                     100
                                  150
                                               200
                                                            NA
  subarctic
Replicated along:
                    || sites ||
body.size:
capybara
                    vampire
             moss
     140
                        190
Replicated along:
                    || species ||
metabolic.rate:
capybara
             moss
                    vampire
      20
                 5
Replicated along:
                    || species ||
REPLICATION DIMENSIONS:
 sites species
      6
              3
```

This function takes three arguments: (1) a list of data frames, (2) a character vector, dimids, with the names identifying the dimensions of replication (i.e. the names of the columns shared between the tables), and (3) a vector, fill, with one element for each data frame giving the value with which to fill in any structural missing values. This last argument is particularly interesting, because we can fill missing abundances with zeros because those site-species combinations were not observed and NA values for the other tables. This data list can now be used in analyses.

It is quite possible that your data are not stored in data frames with columns for both dimensions of replication and variables; this is not a large concern, as the multitable package offers many ways to get your data into a data list (see vignette ????). However, we recommend that researchers at least consider what their data might look like in this format, because we believe that this format best illustrates concepts that will help make multiple-table data easier to understand, manage, and analyze. I now finish with some further elaboration of these concepts.

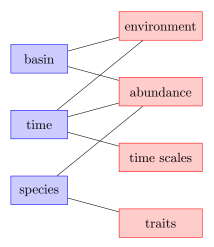


Figure 6: Bipartite graph of the Lac Croche data in Fig. 2.

8 Multiple-table concepts

Shared columns (i.e. dimensions of replication) between tables are expanded by dlcast into the dimensions of the arrays that are used to store each of the other columns (i.e. variables). For example, because the abundance table has two dimensions of replication, it is stored as a two-dimensional matrix in the resulting data frame; whereas, the environment and trait tables have only two dimensions of replication and so are stored in one-dimensional vectors.

The benefits of the distinction between dimensions of replication and variables, is that it provides a common framework for understanding both simple and more complex multiple-table data structures. In particular, the framework allows us to visualize the structure of complex data; for example the Lac Croche zooplankton community data (Fig. 2) has a structure given by Fig. 6. To store these data in a format amenable to dlcast, we would create one data frame for each of the groups of variables (red boxes) and add a column for each dimension of replication (blue boxes) associated with those variables.

With our data in this form, we can easily state the two requirements for using data.list objects: (1) every table must share at least one dimension of replication with at least one other table and (2) at least one table must be replicated along all of the dimensions present in the data set. The first criterion ensures that the tables will relate to each other; the second criterion ensures that some variables will be relatable to all other variables, a property that we feel is necessary for a response variable. We also see that we only need more than one table if some variables are not replicated along all of the dimensions.

9 Conclusion

The structure of data.list objects is sufficiently rich to give rise to a much wider variety of uses than I could cover here. However, this vignette was only intended to illustrate the basic features and concepts of the multitable package, and to justify its utility. My long-term goal with the multitable project in general is to make standard analyses in R (and beyond?) simpler to conduct on complex multiple-table data.

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