

# Working with communities (0.1-639)

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## 1 Introduction

The core of the package is a flexible, extendable representation of an ecological community, described in this vignette. Cheddar's system for plotting communities and statistical analysis of communities is covered in the 'PlotsAndStats' vignette. The 'ImportExport' vignette covers getting community data in to and out of Cheddar. If you are working with collections, for example to see how community structure changes through time, across environmental gradients or resulting from experimental manipulation, read the 'Collections' vignette.

## 2 Datasets

Cheddar contains several published empirical food-web datasets (Table 1).

Community	Notes	References
Benguela	Crude estimate of $M$ ; trophic links have diet fraction	Yodzis (1998)
BroadstoneStream	$M$ and $N$ ; nodes are well resolved	Woodward et al. (2005)
TL84 and TL86	$M$ and $N$ ; nodes are well resolved	Carpenter and Kitchell (1996) Cohen et al. (2003) Jonsson et al. (2005)
SkipwithPond	No $M$ or $N$ ; trophic links have ‘link.evidence’ and ‘link.life.stage’ properties	Warren (1989)
YthanEstuary	$M$ and $N$ for all nodes except detritus; nodes are well-resolved at high trophic levels but poorly resolved at low trophic levels	Hall and Raffaelli (1991) Emmerson and Raffaelli (2004)

Table 1: Community data in Cheddar.  $M$ : body mass.  $N$ : numerical abundance.

## 3 Community representation

A Cheddar community has three aspects:

- *community properties* such as sampling date, lat & long, altitude, temperature and pH,
- *nodes*, which are the names of species together with any associated properties such as mean body mass,  $M$ , and mean numerical abundance,  $N$ , and taxonomic classification,
- the *food web*, defined as the names of each resource-consumer node pair, together with properties such as evidence for the link (e.g. empirically observed or inferred from literature).

The final aspect is optional - Cheddar communities need not contain trophic links. The `LoadCommunity` and `SaveCommunity` functions provide a standard data format, with each aspect stored in its own CSV (Comma-Separated Value) file, described further in the ‘ImportExport’ vignette. Cheddar allows user-defined data to be added to any of these three aspects simply by adding the data to the relevant CSV file. Any data so added will be available to Cheddar’s plotting and analysis functions. Each aspect is accessed using the functions `CPS` (for `Community PropertieS`), `NPS` (for `Node PropertieS`) and `TLPS` (for `Trophic Link PropertieS`) (Table 2). Each of the three community aspects are discussed below. The following examples use the `TL84` dataset, which is from Tuesday Lake in Michigan, USA, sampled in 1984 (Carpenter and Kitchell, 1996; Cohen et al., 2003; Jonsson et al., 2005). The data contain estimates of body mass,  $M$ , and numerical abundance,  $N$ , for each species.

```
> data(TL84) # Load the dataset  
> print(TL84) # A description of the data
```

Tuesday Lake sampled in 1984 containing 56 nodes and 269 trophic links

Aspect	Accessor function	PlotFunction	CSV file
Properties	CPS	n/a	properties.csv
Nodes	NPS	PlotNPS	nodes.csv
Food web	TLPS	PlotTLPS	trophic.links.csv

Table 2: Aspects of a Cheddar community

### 3.1 Community properties

The `CommunityPropertyNames` function returns the names of the community properties.

```
> CommunityPropertyNames(TL84)
```

```
[1] "title"    "M.units"  "N.units"  "lat"       "long"      "habitat"
```

'title' is the only property that a community is guaranteed to contain. The function `CPS` (for **C**ommunity **P**ropertieS) returns a `list` of values.

```
> CPS(TL84)
```

```
$title
```

```
[1] "Tuesday Lake sampled in 1984"
```

```
$M.units
```

```
[1] "kg"
```

```
$N.units
```

```
[1] "m^-3"
```

```
$lat
```

```
[1] 46.21667
```

```
$long
```

```
[1] 89.53333
```

```
$habitat
```

```
[1] "Freshwater pelagic"
```

This shows the latitude and longitude of the lake and tells us the units for body mass,  $M$ , and numerical abundance,  $N$ , are kg and individuals per metre cubed, respectively. Many of the provided communities (Table 1) contain lat, long and habitat. Some communities have more properties. `CPS` lets you get a subset of community properties. For example, to see only the lat and long.

```
> CPS(TL84, c('lat', 'long'))
```

```
$lat
```

```
[1] 46.21667
```

```
$long
```

```
[1] 89.53333
```

CPS also accepts the names of functions that compute community properties. Two such functions are `NumberOfNodes` and `NumberOfTrophicLinks`.

```
> NumberOfNodes(TL84)
[1] 56
> NumberOfTrophicLinks(TL84)
[1] 269
> # A list containing lat, long, number of nodes and number of trophic links
> CPS(TL84, c('lat', 'long', 'NumberOfNodes', 'NumberOfTrophicLinks'))
$lat
[1] 46.21667
$long
[1] 89.53333
$NumberOfNodes
[1] 56
$NumberOfTrophicLinks
[1] 269
```

A named vector can be used to rename values.

```
> CPS(TL84, c('lat', 'long', S='NumberOfNodes', L='NumberOfTrophicLinks'))
$lat
[1] 46.21667
$long
[1] 89.53333
$S
[1] 56
$L
[1] 269
```

Names that are neither properties of the community nor function names result in NA.

```
> # Returns a list containing 'not a property'=NA
> CPS(TL84, c('not a property'))
$`not a property`
[1] NA
```

The related function `CollectionCPS` will be of interest if you are examining collections of communities, described in the ‘Collections’ vignette.

### 3.2 Nodes

Let's use two more Cheddar functions to get some more information about TL84's nodes.

```
> NumberOfNodes(TL84)
```

```
[1] 56
```

```
> NodePropertyNames(TL84)
```

```
[1] "node"          "category"       "M"           "N"           "kingdom"        "phylum"  
[7] "class"         "order"          "family"        "genus"        "species"        "resolved.to"
```

The data contains 56 nodes and `NodePropertyNames` tells us that TL84 contains a lot of information about each node. We can get a table of these node properties by using the `NPS` function. To avoid printing the full table of 56 rows, the examples below use R's `head` and `tail` functions to show just the first or last six rows of the `data.frame` returned by `NPS`.

```
> head(NPS(TL84))
```

	node	category	M	N	kingdom
Nostoc sp.		Nostoc sp.	producer	7.97e-13	2.0e+06
Arthrodesmus sp.		Arthrodesmus sp.	producer	1.52e-12	4.9e+07
Asterionella formosa		Asterionella formosa	producer	1.12e-12	5.0e+06
Cryptomonas sp. 1		Cryptomonas sp. 1	producer	2.03e-13	6.4e+07
Cryptomonas sp. 2		Cryptomonas sp. 2	producer	1.51e-12	2.8e+07
Chroococcus dispersus		Chroococcus dispersus	producer	2.39e-13	2.0e+07
	phylum	class	order	family	
Nostoc sp.	Cyanobacteria	Cyanophyceae	Nostocales	Nostocaceae	
Arthrodesmus sp.	Charophyta	Zygnematophyceae	Desmidiales	Desmidiaceae	
Asterionella formosa	Ochrophyta	Bacillariophyceae	Fragilariales	Fragilariaceae	
Cryptomonas sp. 1	Cryptophyta	Cryptophyceae	Cryptomonadales	Cryptomonadaceae	
Cryptomonas sp. 2	Cryptophyta	Cryptophyceae	Cryptomonadales	Cryptomonadaceae	
Chroococcus dispersus	Cyanobacteria	Cyanophyceae	Chroococcales	Chroococcaceae	
	genus	species	resolved.to		
Nostoc sp.	Nostoc		Species		
Arthrodesmus sp.	Arthrodesmus		Species		
Asterionella formosa	Asterionella	formosa	Species		
Cryptomonas sp. 1	Cryptomonas		Species		
Cryptomonas sp. 2	Cryptomonas		Species		
Chroococcus dispersus	Chroococcus	dispersus	Species		

'node' is the only node property that a community is guaranteed to contain. Many of Cheddar's plotting an analysis functions make use of the 'category' node property; this property is optional but, if included in a community, it should contain one of 'producer', 'invertebrate', 'vert.ecto', 'vert.endo' or should be empty.

```
> # Just body mass  
> head(NPS(TL84, 'M'))
```

```

M
Nostoc sp.      7.97e-13
Arthrodесmus sp. 1.52e-12
Asterionella formosa 1.12e-12
Cryptomonas sp. 1 2.03e-13
Cryptomonas sp. 2 1.51e-12
Chroococcus dispersus 2.39e-13

```

```

> # Body mass and numerical abundance.
> head(NPS(TL84, c('M', 'N')))

```

	M	N
Nostoc sp.	7.97e-13	2.0e+06
Arthrodесmus sp.	1.52e-12	4.9e+07
Asterionella formosa	1.12e-12	5.0e+06
Cryptomonas sp. 1	2.03e-13	6.4e+07
Cryptomonas sp. 2	1.51e-12	2.8e+07
Chroococcus dispersus	2.39e-13	2.0e+07

In addition to first-class node properties like  $M$  and  $N$ , you can also use `NPS` to assemble computed node properties by passing in the name(s) of function(s) that take a community object as the only parameter and return either a vector of length `NumberOfNodes` or a `matrix` or `data.frame` with `NumberOfNodes` rows. Cheddar contains many suitable functions and you can also write your own. For example, it is common to  $\log_{10}$ -transformation  $M$  and  $N$ , which we can do using the `Log10M` and `Log10N` functions.

```
> tail(NPS(TL84, c('Log10M', 'Log10N')))
```

	Log10M	Log10N
Trichocerca cylindrica	-9.420216	4.9116902
Tropocyclops prasinus	-8.164309	4.6919651
Chaoborus punctipennis	-6.522879	4.0791812
Phoxinus eos	-2.995679	0.2944662
Phoxinus neogaeus	-2.931814	-0.8761484
Umbra limi	-2.889410	-0.8794261

You can provide a mix of property and function names.

```
> tail(NPS(TL84, c('Log10M', 'Log10N', 'category', 'phylum')))
```

	Log10M	Log10N	category	phylum
Trichocerca cylindrica	-9.420216	4.9116902	invertebrate	Rotifera
Tropocyclops prasinus	-8.164309	4.6919651	invertebrate	Arthropoda
Chaoborus punctipennis	-6.522879	4.0791812	invertebrate	Arthropoda
Phoxinus eos	-2.995679	0.2944662	vert.ecto	Chordata
Phoxinus neogaeus	-2.931814	-0.8761484	vert.ecto	Chordata
Umbra limi	-2.889410	-0.8794261	vert.ecto	Chordata

The `Log10MNBiomass` function returns a `matrix` of  $\log_{10}$ -transformed body mass,  $M$ , numerical abundance,  $N$ , and biomass,  $B$ , and is a convenient way to get all three of these properties in to a table.

```
> tail(NPS(TL84, c('Log10MNBiomass')))
```

	Log10M	Log10N	Log10Biomass
Trichocerca cylindrica	-9.420216	4.9116902	-4.508526
Tropocyclops prasinus	-8.164309	4.6919651	-3.472344
Chaoborus punctipennis	-6.522879	4.0791812	-2.443697
Phoxinus eos	-2.995679	0.2944662	-2.701212
Phoxinus neogaeus	-2.931814	-0.8761484	-3.807962
Umbra limi	-2.889410	-0.8794261	-3.768836

We can use NPS to assemble a table showing node degree: the number of trophic links in to and out of that node. Cheddar contains three functions that compute a different aspect of node degree.

```
> nps <- NPS(TL84, c('InDegree', 'OutDegree', 'Degree'))
> head(nps)
```

	InDegree	OutDegree	Degree
Nostoc sp.	0	4	4
Arthrodemesmus sp.	0	3	3
Asterionella formosa	0	0	0
Cryptomonas sp. 1	0	18	18
Cryptomonas sp. 2	0	5	5
Chroococcus dispersus	0	18	18

```
> # This is always true for all nodes
> all(nps$Degree == nps$InDegree+nps$OutDegree)
```

```
[1] TRUE
```

Some readers will be more familiar with the terms ‘trophic vulnerability’ and ‘trophic generality’; the functions `TrophicVulnerability` and `TrophicGenerality` are synonyms for `OutDegree` and `InDegree` respectively. Cannibalistic links count twice towards `Degree` - one link going out and one going in. The cannibalistic fish *Umbra limi* in TL84 has no consumers other than itself so it has an `OutDegree` of one.

```
> IsCannibal(TL84)[‘Umbra limi’]
```

```
Umbra limi
TRUE
```

```
> InDegree(TL84)[“Umbra limi”]
```

```
Umbra limi
12
```

```
> OutDegree(TL84)[“Umbra limi”]
```

```
Umbra limi
1
```

```
> Degree(TL84)[“Umbra limi”]
```

```
Umbra limi
13
```

We can combine some of these functions to investigate allometric degree distribution (Jonsson et al., 2005; Otto et al., 2007; Digel et al., 2011; Jacob et al., 2011), which describe how species' numbers of trophic links scale with their log-transformed body masses.

```
> tail(NPS(TL84, c('Log10M', 'OutDegree', 'InDegree', 'Degree')))
```

	Log10M	OutDegree	InDegree	Degree
Trichocerca cylindrica	-9.420216	4	6	10
Tropocyclops prasinus	-8.164309	7	16	23
Chaoborus punctipennis	-6.522879	4	22	26
Phoxinus eos	-2.995679	1	9	10
Phoxinus neogaeus	-2.931814	1	9	10
Umbra limi	-2.889410	1	12	13

Some authors have been interested in how trophic level varies with body mass (Jacob et al., 2011). Two more functions suitable for use with NPS are `PreyAveragedTrophicLevel` and `ChainAveragedTrophicLevel`, which give different measures of each node's trophic level in the food web; these two functions, and others related to trophic level, are discussed further in section 3.3.

```
> tail(NPS(TL84, c('Log10M', 'PreyAveragedTrophicLevel',
  'ChainAveragedTrophicLevel')))
```

	Log10M	PreyAveragedTrophicLevel	ChainAveragedTrophicLevel
Trichocerca cylindrica	-9.420216	2.000000	2.000000
Tropocyclops prasinus	-8.164309	3.142857	3.333333
Chaoborus punctipennis	-6.522879	3.171344	4.602527
Phoxinus eos	-2.995679	3.529951	5.168337
Phoxinus neogaeus	-2.931814	3.529951	5.168337
Umbra limi	-2.889410	3.802678	5.835003

The column titles for the trophic-level measures are very long. We can use a named vector to get shortened column titles.

```
> tail(NPS(TL84, c('Log10M', PATL='PreyAveragedTrophicLevel',
  CATL='ChainAveragedTrophicLevel')))
```

	Log10M	PATL	CATL
Trichocerca cylindrica	-9.420216	2.000000	2.000000
Tropocyclops prasinus	-8.164309	3.142857	3.333333
Chaoborus punctipennis	-6.522879	3.171344	4.602527
Phoxinus eos	-2.995679	3.529951	5.168337
Phoxinus neogaeus	-2.931814	3.529951	5.168337
Umbra limi	-2.889410	3.802678	5.835003

NPS also allows parameters to be passed to functions. This is demonstrated using the `TrophicSpecies` function: in order to account for different levels of taxonomic resolution and other biases, researchers often lump biological species together. Species in the food web that have the same resources and consumers are the same 'trophic species' (Briand and Cohen, 1984; Pimm et al., 1991; Williams and Martinez, 2000). The `TrophicSpecies` function computes these IDs.

```
> tail(TrophicSpecies(TL84))
```

Trichocerca cylindrica	Tropocyclops prasinus	Chaoborus punctipennis	
16	13	20	
Phoxinus eos	Phoxinus neogaeus	Umbra limi	
21	21	22	

Some analyses (e.g. Jonsson et al., 2005) exclude isolated species when computing trophic species numbers. Isolated species are those nodes that consume no others and have no consumers (Section 3.3.4). To compare the effect of including or excluding isolated species we can pass the function to NPS twice, once setting the ‘include.isolated’ parameter.

```
> head(NPS(TL84, list(TS.iso='TrophicSpecies',
                      TS.no.iso=list('TrophicSpecies', include.isolated=FALSE))))
```

	TS.iso	TS.no.iso
Nostoc sp.	1	1
Arthrodesmus sp.	2	2
Asterionella formosa	3	NA
Cryptomonas sp. 1	4	3
Cryptomonas sp. 2	5	4
Chroococcus dispersus	4	3

*Asterionella formosa* is an isolated species so has been given a trophic species of NA in the ‘TS.no.iso’ column. The **LumpTrophicSpecies** function lumps nodes together using these IDs (Section 4.5). The second argument to NPS can therefore be defined as a list containing either names of first class properties, names of functions that take only a community or lists in which the first element is the name of a function that takes a community and subsequent elements are *named* arguments to that function. Names of the list are column names in the returned **data.frame**.

NPS therefore makes it very easy to assemble tables of properties either for analysis or for presentation in a manuscript. The example below recreates the first ten rows of Jonsson et al. (2005), Appendix 1A (p74–75).

```
> head(NPS(TL84, list('category', BM='M', 'NA'='N',
                      TS=list('TrophicSpecies', include.isolated=FALSE),
                      TH=list('TrophicHeight', include.isolated=FALSE))),  
10)
```

	category	BM	NA	TS	TH
Nostoc sp.	producer	7.97e-13	2.0e+06	1	1
Arthrodesmus sp.	producer	1.52e-12	4.9e+07	2	1
Asterionella formosa	producer	1.12e-12	5.0e+06	NA	NA
Cryptomonas sp. 1	producer	2.03e-13	6.4e+07	3	1
Cryptomonas sp. 2	producer	1.51e-12	2.8e+07	4	1
Chroococcus dispersus	producer	2.39e-13	2.0e+07	3	1
Closteriopsis longissimus	producer	2.37e-13	1.0e+08	5	1
Chrysosphaerella longispina	producer	8.31e-10	4.0e+06	NA	NA
Dinobryon bavaricum	producer	2.44e-12	3.0e+07	6	1
Dinobryon cylindricum	producer	1.57e-12	3.0e+06	1	1

Some values in this table are different to those presented by Jonsson et al. (2005) in their Appendix 1A. Firstly, the numerical abundance values for zooplankton are different. Values in their table "...should

be multiplied by 6 to convert them to concentrations in the epilimnion" (Jonsson et al., 2005), and our data incorporate that conversion. Secondly, the values of trophic height presented are slightly different for species at higher trophic levels because of the different methods used to break cycles (see the help for Cheddar's **TrophicSpecies** function). It is not clear from the text of Jonsson et al. (2005) exactly how they broke cycles. Because Cheddar is open source, users can refer readers to the function and version used to completely specify the algorithm used.

**NPS** returns **NA** for any names that are neither a first-class property of the community nor the name of a function.

```
> head(NPS(TL84, c('Not a property or function')))
```

Not a property or function	
Nostoc sp.	NA
Arthrodemus sp.	NA
Asterionella formosa	NA
Cryptomonas sp. 1	NA
Cryptomonas sp. 2	NA
Chroococcus dispersus	NA

### 3.3 Food web

**NumberOfTrophicLinks** returns the number of trophic links that the community contains.

```
> NumberOfTrophicLinks(TL84)
```

```
[1] 269
```

Cheddar communities need not contain trophic links so this function might return zero. The following sections describe some different ways in which to view and analyse food webs in cheddar.

#### 3.3.1 Resource-consumer pairs

**TLPS** (for **Trophic Link PropertieS**) returns a **data.frame** of trophic links pairs (or **NULL** if the community has no food web). The **data.frame** always contains the columns 'resource' and 'consumer'.

```
> head(TLPS(TL84))
```

	resource	consumer
1	Cryptomonas sp. 1	Ascomorpha eucadis
2	Chroococcus dispersus	Ascomorpha eucadis
3	Unclassified flagellates	Ascomorpha eucadis
4	Chromulina sp.	Ascomorpha eucadis
5	Selenastrum minutum	Ascomorpha eucadis
6	Trachelomonas sp.	Ascomorpha eucadis

**TLPS** takes a parameter **node.properties**, which should be a vector of names suitable for passing to **NPS**. You can therefore use functions and names and can pass parameters to functions, just as in the **NPS** examples above.

```
> head(TLPS(TL84, node.properties='M'))
```

	resource	consumer	resource.M
Cryptomonas sp. 1	Cryptomonas sp. 1	Ascomorpha eucadis	2.03e-13
Chroococcus dispersus	Chroococcus dispersus	Ascomorpha eucadis	2.39e-13
Unclassified flagellates	Unclassified flagellates	Ascomorpha eucadis	3.46e-13
Chromulina sp.	Chromulina sp.	Ascomorpha eucadis	3.03e-14
Selenastrum minutum	Selenastrum minutum	Ascomorpha eucadis	2.72e-13
Trachelomonas sp.	Trachelomonas sp.	Ascomorpha eucadis	1.75e-13
	consumer.M		
Cryptomonas sp. 1	1.4e-10		
Chroococcus dispersus	1.4e-10		
Unclassified flagellates	1.4e-10		
Chromulina sp.	1.4e-10		
Selenastrum minutum	1.4e-10		
Trachelomonas sp.	1.4e-10		

```
> head(TLPS(TL84, node.properties=c('M', 'Biomass')))
```

	resource	consumer	resource.M
Cryptomonas sp. 1	Cryptomonas sp. 1	Ascomorpha eucadis	2.03e-13
Chroococcus dispersus	Chroococcus dispersus	Ascomorpha eucadis	2.39e-13
Unclassified flagellates	Unclassified flagellates	Ascomorpha eucadis	3.46e-13
Chromulina sp.	Chromulina sp.	Ascomorpha eucadis	3.03e-14
Selenastrum minutum	Selenastrum minutum	Ascomorpha eucadis	2.72e-13
Trachelomonas sp.	Trachelomonas sp.	Ascomorpha eucadis	1.75e-13
	resource.Biomass	consumer.M	consumer.Biomass
Cryptomonas sp. 1	1.2992e-05	1.4e-10	1.932e-06
Chroococcus dispersus	4.7800e-06	1.4e-10	1.932e-06
Unclassified flagellates	6.5048e-04	1.4e-10	1.932e-06
Chromulina sp.	4.5147e-06	1.4e-10	1.932e-06
Selenastrum minutum	5.4400e-05	1.4e-10	1.932e-06
Trachelomonas sp.	3.8850e-05	1.4e-10	1.932e-06

```
> head(TLPS(TL84, node.properties=c('M', B='Biomass')))
```

	resource	consumer	resource.M
Cryptomonas sp. 1	Cryptomonas sp. 1	Ascomorpha eucadis	2.03e-13
Chroococcus dispersus	Chroococcus dispersus	Ascomorpha eucadis	2.39e-13
Unclassified flagellates	Unclassified flagellates	Ascomorpha eucadis	3.46e-13
Chromulina sp.	Chromulina sp.	Ascomorpha eucadis	3.03e-14
Selenastrum minutum	Selenastrum minutum	Ascomorpha eucadis	2.72e-13
Trachelomonas sp.	Trachelomonas sp.	Ascomorpha eucadis	1.75e-13
	resource.B	consumer.M	consumer.B
Cryptomonas sp. 1	1.2992e-05	1.4e-10	1.932e-06
Chroococcus dispersus	4.7800e-06	1.4e-10	1.932e-06
Unclassified flagellates	6.5048e-04	1.4e-10	1.932e-06
Chromulina sp.	4.5147e-06	1.4e-10	1.932e-06
Selenastrum minutum	5.4400e-05	1.4e-10	1.932e-06
Trachelomonas sp.	3.8850e-05	1.4e-10	1.932e-06

`TLPS` takes a parameter `link.properties`, which should be a vector of names that are either first-class trophic-link properties or are functions. Functions should take a community as the first parameter and a second parameter that is a `data.frame` containing the columns ‘resource’ and ‘consumer’. They should return either a vector of length `NumberOfTrophicLinks` or a matrix or `data.frame` with `NumberOfTrophicLinks` rows.

### 3.3.2 Trophic-link properties

Food web data in Cheddar communities can be augmented with extra information. The dataset of `SkipwithPond` (Warren, 1989) contains two such properties: ‘link.evidence’ and ‘link.life.stage’.

```
> data(SkipwithPond)
> head(TLPS(SkipwithPond))
```

		resource	consumer	link.evidence
1	Small oligochaetes (principally Enchytraeidae)	Polycelis tenuis		Inferred
2		Lumbriculus variegatus	Polycelis tenuis	Inferred
3		Procladius sagittalis	Polycelis tenuis	Inferred
4		Corynoneura scutellata	Polycelis tenuis	Inferred
5		Chironomus dorsalis	Polycelis tenuis	Known
6		Glyptotendipes pallens	Polycelis tenuis	Known
		link.life.stage		
1	All life stages			
2	All life stages			
3	All life stages			
4	All life stages			
5	All life stages			
6	All life stages			

`TrophicLinkPropertyNames` returns the names of the trophic-link properties in a community.

```
> TrophicLinkPropertyNames(SkipwithPond)
```

```
[1] "resource"      "consumer"       "link.evidence"   "link.life.stage"
```

`TLPS` accepts a ‘link.properties’ parameter. You can use this to get a subset of the first-class link properties. The columns ‘resource’ and ‘consumer’ are always returned.

```
> head(TLPS(SkipwithPond, link.properties='link.evidence'))
```

		resource	consumer	link.evidence
1	Small oligochaetes (principally Enchytraeidae)	Polycelis tenuis		Inferred
2		Lumbriculus variegatus	Polycelis tenuis	Inferred
3		Procladius sagittalis	Polycelis tenuis	Inferred
4		Corynoneura scutellata	Polycelis tenuis	Inferred
5		Chironomus dorsalis	Polycelis tenuis	Known
6		Glyptotendipes pallens	Polycelis tenuis	Known

`TLPS` takes a parameter `link.properties`, which should be a vector of names that are either first-class trophic-link properties or are functions. Functions should take a community as the only parameter.

They should return either a vector of length `NumberOfTrophicLinks` or a matrix or data.frame with `NumberOfTrophicLinks` rows. This is illustrated by the code fragment below, which uses the `Log10RCMRatio` function to get the  $\log_{10}$ -transformed ratio between body mass of the resource and consumer in each trophic link in TL84.

```
> head(TLPS(TL84, link.properties='Log10RCMRatio'))
```

	resource	consumer	Log10RCMRatio
1	Cryptomonas sp. 1	Ascomorpha eucadis	-2.838632
2	Chroococcus dispersus	Ascomorpha eucadis	-2.767730
3	Unclassified flagellates	Ascomorpha eucadis	-2.607052
4	Chromulina sp.	Ascomorpha eucadis	-3.664685
5	Selenastrum minutum	Ascomorpha eucadis	-2.711559
6	Trachelomonas sp.	Ascomorpha eucadis	-2.903090

You can combine `node.properties` and `link.properties`.

```
> head(TLPS(TL84, node.properties='Log10M', link.properties='Log10RCMRatio'))
```

	resource	consumer	Log10RCMRatio	resource.Log10M	consumer.Log10M
1	Cryptomonas sp. 1	Ascomorpha eucadis	-2.838632	-12.69250	-9.853872
2	Chroococcus dispersus	Ascomorpha eucadis	-2.767730	-12.62160	-9.853872
3	Unclassified flagellates	Ascomorpha eucadis	-2.607052	-12.46092	-9.853872
4	Chromulina sp.	Ascomorpha eucadis	-3.664685	-13.51856	-9.853872
5	Selenastrum minutum	Ascomorpha eucadis	-2.711559	-12.56543	-9.853872
6	Trachelomonas sp.	Ascomorpha eucadis	-2.903090	-12.75696	-9.853872

### 3.3.3 Predation matrix

The `PredationMatrix` function returns an R `matrix` object. The matrix returned by the code fragment below is 56 x 56 and so is not shown for brevity.

```
> pm <- PredationMatrix(TL84)
```

In the example above, all entries in ‘pm’ are either 0 or 1. This summation below computes the number of 1s in the matrix, which is the same as the number of trophic links in the community.

```
> sum(pm)
```

```
[1] 269
```

```
> NumberOfTrophicLinks(TL84)
```

```
[1] 269
```

Data that contain estimates of link strength can be used to construct a weighted predation matrix, such as the `Benguela` dataset, which contains the ‘`diet.fraction`’ node property (Yodzis, 1998).

```
> data(Benguela)
> pm <- PredationMatrix(Benguela, weight='diet.fraction')
```

More information about link strengths is in Section 3.3.8.

### 3.3.4 Node connectivity

A node in a community can be defined as falling in to one of four categories (Table 3). A node will

Category	Description
Isolated	No resources or consumers
Basal	No resources and one or more consumers
Top-level	One or more resources and no consumers
Intermediate	Nodes not fitting any of the above categories

Table 3: Node connectivity. Cannibalistic links are disregarded.

satisfy only one of the above four definitions. These definitions allow three additional definitions (Table 4). For each of the seven definitions (Tables 3 and 4), ‘X’, there are three functions: `IsXNode`, `XNodes`

Category	Description
Connected	Basal, Intermediate or Top-level
Non-basal	Isolated, Intermediate or Top-level
Non-top-level	Isolated, Basal or Intermediate

Table 4: Additional node connectivity

and `FractionXNodes`. The first returns a vector of type `logical` of length `NumberOfNodes`; values are `TRUE` for nodes that fit the definition of ‘X’. The second returns the names of nodes for which `IsXNode` returns `TRUE`. The third returns the proportion of nodes in the community that fit the definition of ‘X’. For example, a community’s isolated species can be accessed by using `IsolatedNodes`.

```
> IsolatedNodes(TL84)

[1] "Asterionella formosa"           "Chrysosphaerella longispina"
[3] "Diceras sp."                  "Rhizosolenia sp."
[5] "Spinocosmarium sp."           "Staurastrum sp."
```

We can use the `IsXNode` functions together with `NPS` to see a table of connectivity for the whole community.

```
> connectivity <- NPS(TL84, c(Basal='IsBasalNode',
                                Isolated='IsIsolatedNode',
                                Intermediate='IsIntermediateNode',
                                TopLevel='IsTopLevelNode'))
> connectivity
```

	Basal	Isolated	Intermediate	TopLevel
<i>Nostoc</i> sp.	TRUE	FALSE	FALSE	FALSE
<i>Arthrodesmus</i> sp.	TRUE	FALSE	FALSE	FALSE
<i>Asterionella formosa</i>	FALSE	TRUE	FALSE	FALSE
<i>Cryptomonas</i> sp. 1	TRUE	FALSE	FALSE	FALSE
<i>Cryptomonas</i> sp. 2	TRUE	FALSE	FALSE	FALSE
<i>Chroococcus dispersus</i>	TRUE	FALSE	FALSE	FALSE
<i>Closteriopsis longissimus</i>	TRUE	FALSE	FALSE	FALSE
<i>Chrysosphaerella longispina</i>	FALSE	TRUE	FALSE	FALSE
<i>Dinobryon bavaricum</i>	TRUE	FALSE	FALSE	FALSE
<i>Dinobryon cylindricum</i>	TRUE	FALSE	FALSE	FALSE
<i>Dactylococcopsis fascicularis</i>	TRUE	FALSE	FALSE	FALSE
<i>Diceras</i> sp.	FALSE	TRUE	FALSE	FALSE
<i>Dictyosphaerium pulchellum</i>	TRUE	FALSE	FALSE	FALSE
<i>Dinobryon sertularia</i>	TRUE	FALSE	FALSE	FALSE
<i>Dinobryon sociale</i>	TRUE	FALSE	FALSE	FALSE
<i>Glenodinium quadridens</i>	TRUE	FALSE	FALSE	FALSE
<i>Microcystis aeruginosa</i>	TRUE	FALSE	FALSE	FALSE
<i>Mallomonas</i> sp. 1	TRUE	FALSE	FALSE	FALSE
<i>Mallomonas</i> sp. 2	TRUE	FALSE	FALSE	FALSE
Unclassified flagellates	TRUE	FALSE	FALSE	FALSE
<i>Peridinium limbatum</i>	TRUE	FALSE	FALSE	FALSE
<i>Peridinium cinctum</i>	TRUE	FALSE	FALSE	FALSE
<i>Peridinium pulsillum</i>	TRUE	FALSE	FALSE	FALSE
<i>Peridinium wisconsinense</i>	TRUE	FALSE	FALSE	FALSE
<i>Chromulina</i> sp.	TRUE	FALSE	FALSE	FALSE
<i>Rhizosolenia</i> sp.	FALSE	TRUE	FALSE	FALSE
<i>Selenastrum minutum</i>	TRUE	FALSE	FALSE	FALSE
<i>Spinocosmarium</i> sp.	FALSE	TRUE	FALSE	FALSE
<i>Staurastrum</i> sp.	FALSE	TRUE	FALSE	FALSE
<i>Synedra</i> sp.	TRUE	FALSE	FALSE	FALSE
<i>Trachelomonas</i> sp.	TRUE	FALSE	FALSE	FALSE
<i>Ascomorpha eucadis</i>	FALSE	FALSE	TRUE	FALSE
<i>Synchaeta</i> sp.	FALSE	FALSE	TRUE	FALSE
<i>Bosmina longirostris</i>	FALSE	FALSE	TRUE	FALSE
<i>Conochilus</i> (solitary)	FALSE	FALSE	TRUE	FALSE
<i>Cyclops varians rubellus</i>	FALSE	FALSE	TRUE	FALSE
<i>Diaphanosoma leuchtenbergianum</i>	FALSE	FALSE	TRUE	FALSE
<i>Daphnia pulex</i>	FALSE	FALSE	TRUE	FALSE
<i>Filinia longispina</i>	FALSE	FALSE	TRUE	FALSE
<i>Conochiloides dossuarius</i>	FALSE	FALSE	TRUE	FALSE
<i>Gastropus stylifer</i>	FALSE	FALSE	TRUE	FALSE
<i>Holopedium gibberum</i>	FALSE	FALSE	TRUE	FALSE
<i>Kellicottia</i> sp.	FALSE	FALSE	TRUE	FALSE
<i>Keratella cochlearis</i>	FALSE	FALSE	TRUE	FALSE
<i>Keratella testudo</i>	FALSE	FALSE	TRUE	FALSE
<i>Leptodiaptomus siciloides</i>	FALSE	FALSE	TRUE	FALSE
<i>Orthocyclops modestus</i>	FALSE	FALSE	TRUE	FALSE

Ploesoma sp.	FALSE	FALSE	TRUE	FALSE
Polyarthra vulgaris	FALSE	FALSE	TRUE	FALSE
Trichocerca multicrinis	FALSE	FALSE	TRUE	FALSE
Trichocerca cylindrica	FALSE	FALSE	TRUE	FALSE
Tropocyclops prasinus	FALSE	FALSE	TRUE	FALSE
Chaoborus punctipennis	FALSE	FALSE	TRUE	FALSE
Phoxinus eos	FALSE	FALSE	TRUE	FALSE
Phoxinus neogaeus	FALSE	FALSE	TRUE	FALSE
Umbra limi	FALSE	FALSE	FALSE	TRUE

Because nodes can fit only one of the definitions in Table 3, each row in the `connectivity` `data.frame` should have one, and only one, value of TRUE. We can verify this by summing each row using R's `apply` function.

```
> all(1==apply(connectivity, 1, sum))
```

```
[1] TRUE
```

The following summations are also 1.

```
> sum(FractionBasalNodes(TL84),
      FractionIntermediateNodes(TL84),
      FractionTopLevelNodes(TL84),
      FractionIsolatedNodes(TL84))
```

```
[1] 1
```

```
> sum(FractionConnectedNodes(TL84),
      FractionIsolatedNodes(TL84))
```

```
[1] 1
```

```
> sum(FractionBasalNodes(TL84),
      FractionNonBasalNodes(TL84))
```

```
[1] 1
```

```
> sum(FractionTopLevelNodes(TL84),
      FractionNonTopLevelNodes(TL84))
```

```
[1] 1
```

### 3.3.5 Trophic chains

Some network properties and analyses require knowledge of every unique path - 'trophic chain' - through the food-web. A trophic chain starts with a basal node (Section 3.3.4) and ends when it is not possible to add nodes that are not already in the chain. Loops and cannibalism are therefore ignored when computing trophic chains. For communities that have one or more top-level nodes each trophic chain will end with a top-level node. The 'length' of a chain is defined as the number of links that it contains, i.e. the number of nodes in the chain minus one.

Cheddar provides two functions for examining trophic chains. The `TrophicChains` function returns a `data.frame` of trophic chains.

```

> tc <- TrophicChains(TL84)
> dim(tc)

[1] 5988     8

```

There are 5988 unique chains in the food web and the longest chains contain 8 nodes. Let's look at the first 20 chains.

```
> head(tc, 20)
```

	Node.1	Node.2	Node.3	Node.4
1	Nostoc sp.	Diaphanosoma leuchtenbergianum	Umbra limi	
2	Nostoc sp.	Daphnia pulex	Umbra limi	
3	Nostoc sp.	Holopedium gibberum	Umbra limi	
4	Nostoc sp.	Leptodiaptomus siciloides	Umbra limi	
5	Nostoc sp.	Diaphanosoma leuchtenbergianum	Chaoborus punctipennis	Umbra limi
6	Nostoc sp.	Diaphanosoma leuchtenbergianum	Phoxinus eos	Umbra limi
7	Nostoc sp.	Diaphanosoma leuchtenbergianum	Phoxinus neogaeus	Umbra limi
8	Nostoc sp.	Daphnia pulex	Chaoborus punctipennis	Umbra limi
9	Nostoc sp.	Daphnia pulex	Phoxinus eos	Umbra limi
10	Nostoc sp.	Daphnia pulex	Phoxinus neogaeus	Umbra limi
11	Nostoc sp.	Holopedium gibberum	Phoxinus eos	Umbra limi
12	Nostoc sp.	Holopedium gibberum	Phoxinus neogaeus	Umbra limi
13	Nostoc sp.	Leptodiaptomus siciloides	Cyclops varians rubellus	Umbra limi
14	Nostoc sp.	Leptodiaptomus siciloides	Orthocyclops modestus	Umbra limi
15	Nostoc sp.	Leptodiaptomus siciloides	Tropocyclops prasinus	Umbra limi
16	Nostoc sp.	Leptodiaptomus siciloides	Chaoborus punctipennis	Umbra limi
17	Nostoc sp.	Leptodiaptomus siciloides	Phoxinus eos	Umbra limi
18	Nostoc sp.	Leptodiaptomus siciloides	Phoxinus neogaeus	Umbra limi
19	Nostoc sp.	Diaphanosoma leuchtenbergianum	Chaoborus punctipennis	Phoxinus eos
20	Nostoc sp.	Diaphanosoma leuchtenbergianum	Chaoborus punctipennis	Phoxinus neogaeus
	Node.5	Node.6	Node.7	Node.8

```

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17

```

```
18  
19 Umbra limi  
20 Umbra limi
```

Every chain starts with a basal node.

```
> BasalNodes(TL84)
```

```
[1] "Nostoc sp."          "Arthrodесmus sp."  
[3] "Cryptomonas sp. 1"    "Cryptomonas sp. 2"  
[5] "Chroococcus dispersus" "Closteriopsis longissimus"  
[7] "Dinobryon bavaricum"   "Dinobryon cylindricum"  
[9] "Dactylococcopsis fascicularis" "Dictyosphaerium pulchellum"  
[11] "Dinobryon sertularia"   "Dinobryon sociale"  
[13] "Glenodinium quadridens" "Microcystis aeruginosa"  
[15] "Mallomonas sp. 1"       "Mallomonas sp. 2"  
[17] "Unclassified flagellates" "Peridinium limbatum"  
[19] "Peridinium cinctum"      "Peridinium pulsillum"  
[21] "Peridinium wisconsinense" "Chromulina sp."  
[23] "Selenastrum minutum"     "Synedra sp."  
[25] "Trachelomonas sp."
```

```
> # The first node in each chain  
> first <- tc[,1]  
> all(unique(first) %in% BasalNodes(TL84)) # TRUE
```

```
[1] TRUE
```

Tuesday Lake 1984 has a single top-level consumer so every trophic chain ends with this consumer.

```
> TopLevelNodes(TL84)
```

```
[1] "Umbra limi"
```

```
> # The last node in each chain  
> last <- apply(tc, 1, function(row) row[max(which("!="row))])  
> unique(last)
```

```
[1] "Umbra limi"
```

Just as with TLPS, `TrophicChains` accepts a ‘node.properties’ parameter that you can use to add node properties to the returned. For example, to get a table containing the  $\log_{10}$ -transformed body mass of each node in every chain.

```
> tc.with.log10M <- TrophicChains(TL84, node.properties='Log10M')
```

The food web of the Bengula marine ecosystem (Yodzis, 1998) does not have any top-level nodes. All chains start with a basal node (by definition) but, for this community, all chains end with an intermediate node.

```
> TopLevelNodes(Benguela)
```

```

character(0)

> tc <- TrophicChains(Benguela)
> last <- apply(tc, 1, function(row) row[max(which("")!=row)])
> unique(last)

[1] "Sharks" "Seals"  "Birds"

> IsIntermediateNode(Benguela)[unique(last)] 

Sharks  Seals  Birds
TRUE    TRUE   TRUE

```

The second function - `TrophicChainsStats` - returns a list of simple statistics about trophic chains.

```
> chain.stats <- TrophicChainsStats(TL84)
```

The ‘chain.lengths’ item contains the length of every unique food chain.

```
> length(chain.stats$chain.lengths)      # 5,988 chains
```

```
[1] 5988
```

```
> summary(chain.stats$chain.lengths)
```

Min.	1st Qu.	Median	Mean	3rd Qu.	Max.
2.000	4.000	5.000	4.835	6.000	7.000

The ‘node.pos.counts’ item is a matrix of `NumberOfNodes` rows and `1+max(chain.lengths)` columns. Elements are the number of chains in which a node appear in that position.

```
> dim(chain.stats$node.pos.counts)      # 56 nodes. Longest chain contains 8 nodes
```

```
[1] 56  8
```

Basal nodes only have counts in the first column.

```
> chain.stats$node.pos.counts[BasalNodes(TL84), ]
```

	[,1]	[,2]	[,3]	[,4]	[,5]	[,6]	[,7]	[,8]
Nostoc sp.	75	0	0	0	0	0	0	0
Arthrodemesmus sp.	69	0	0	0	0	0	0	0
Cryptomonas sp. 1	840	0	0	0	0	0	0	0
Cryptomonas sp. 2	81	0	0	0	0	0	0	0
Chroococcus dispersus	840	0	0	0	0	0	0	0
Closteriopsis longissimus	120	0	0	0	0	0	0	0
Dinobryon bavaricum	6	0	0	0	0	0	0	0
Dinobryon cylindricum	75	0	0	0	0	0	0	0
Dactylococcopsis fascicularis	75	0	0	0	0	0	0	0
Dictyosphaerium pulchellum	81	0	0	0	0	0	0	0
Dinobryon sertularia	9	0	0	0	0	0	0	0
Dinobryon sociale	81	0	0	0	0	0	0	0

Glenodinium quadridens	84	0	0	0	0	0	0	0
Microcystis aeruginosa	6	0	0	0	0	0	0	0
Mallomonas sp. 1	9	0	0	0	0	0	0	0
Mallomonas sp. 2	69	0	0	0	0	0	0	0
Unclassified flagellates	840	0	0	0	0	0	0	0
Peridinium limbatum	6	0	0	0	0	0	0	0
Peridinium cinctum	9	0	0	0	0	0	0	0
Peridinium pulsillum	81	0	0	0	0	0	0	0
Peridinium wisconsinense	6	0	0	0	0	0	0	0
Chromulina sp.	840	0	0	0	0	0	0	0
Selenastrum minutum	840	0	0	0	0	0	0	0
Synedra sp.	6	0	0	0	0	0	0	0
Trachelomonas sp.	840	0	0	0	0	0	0	0

Consumers only have counts in columns two and higher.

```
> chain.stats$node.pos.counts[c(IntermediateNodes(TL84), TopLevelNodes(TL84)), ]
```

	[,1]	[,2]	[,3]	[,4]	[,5]	[,6]	[,7]	[,8]
Ascomorpha eucadis	0	378	0	0	0	0	0	0
Synchaeta sp.	0	378	0	0	0	0	0	0
Bosmina longirostris	0	66	0	0	0	0	0	0
Conochilus (solitary)	0	399	0	0	0	0	0	0
Cyclops varians rubellus	0	0	2304	1152	0	0	0	0
Diaphanosoma leuchtenbergianum	0	84	0	0	0	0	0	0
Daphnia pulex	0	150	108	0	0	0	0	0
Filinia longispina	0	342	0	0	0	0	0	0
Conochilooides dossuarius	0	399	0	0	0	0	0	0
Gastropus stylifer	0	342	0	0	0	0	0	0
Holopedium gibberum	0	57	0	0	0	0	0	0
Kellicottia sp.	0	342	0	0	0	0	0	0
Keratella cochlearis	0	378	0	0	0	0	0	0
Keratella testudo	0	342	0	0	0	0	0	0
Leptodiaptomus siciloides	0	960	0	0	0	0	0	0
Orthocyclops modestus	0	0	576	1152	1152	0	0	0
Ploesoma sp.	0	342	0	0	0	0	0	0
Polyarthra vulgaris	0	342	0	0	0	0	0	0
Trichocerca multicrinis	0	342	0	0	0	0	0	0
Trichocerca cylindrica	0	342	0	0	0	0	0	0
Tropocyclops prasinus	0	0	2304	1152	0	0	0	0
Chaoborus punctipennis	0	3	438	918	1152	576	0	0
Phoxinus eos	0	0	86	452	690	576	192	0
Phoxinus neogaeus	0	0	86	452	690	576	192	0
Umbra limi	0	0	86	624	1594	1956	1344	384

All counts are zero for isolated nodes.

```
> chain.stats$node.pos.counts[IsolatedNodes(TL84), ]
```

	[,1]	[,2]	[,3]	[,4]	[,5]	[,6]	[,7]	[,8]
Asterionella formosa	0	0	0	0	0	0	0	0
Chrysosphaerella longispina	0	0	0	0	0	0	0	0
Diceras sp.	0	0	0	0	0	0	0	0
Rhizosolenia sp.	0	0	0	0	0	0	0	0
Spinocosmarium sp.	0	0	0	0	0	0	0	0
Staurastrum sp.	0	0	0	0	0	0	0	0

If your analysis requires only simple statistics about trophic chains, the `TrophicChainsStats` function is more suitable than `TrophicChains` because it is faster and requires less memory.

```
> system.time(tc <- TrophicChains(TL84))
  user  system elapsed
 0.013   0.000   0.013

> system.time(stats <- TrophicChainsStats(TL84))
  user  system elapsed
 0.012   0.000   0.012
```

The difference in speed will be greater for communities that contain a large number of nodes and trophic chains such as the Skipwith Pond dataset, which has more than  $10^5$  unique chains.

### 3.3.6 Large numbers of trophic chains

The number of possible trophic chains rapidly increases with the number of nodes and links. Let's examine the relationship between number of trophic chains and number of nodes in communities in which every consumer eats every other node.

```
> HighlyConnected <- function(n)
{
  # Returns a community containing a single producer and n consumers, all
  # of whom eat everything else
  consumers <- paste('Consumer', 1:n)
  tl <- data.frame(resource=c(rep('Producer', n), rep(consumers, each=n)),
                    consumer=consumers)
  return (Community(nodes=data.frame(node=c('Producer', consumers)),
                     trophic.links=tl,
                     properties=list(title='test')))
}
> # A list of communities of between 1 and 8 consumers
> n <- 8
> communities <- lapply(1:n, HighlyConnected)
> # A list of statistics about each community
> stats <- lapply(communities, TrophicChainsStats)
> # Extract the chain lengths
> cl <- lapply(stats, '[[', 'chain.lengths')
> # The number of chains
> n.chains <- sapply(cl, length)
> # The number of chains in each community
> cbind(n.consumers=1:n, longest.chain=sapply(cl, max), n.chains=n.chains)
```

	n.consumers	longest.chain	n.chains
[1,]	1	1	1
[2,]	2	2	2
[3,]	3	3	6
[4,]	4	4	24
[5,]	5	5	120
[6,]	6	6	720
[7,]	7	7	5040
[8,]	8	8	40320

So with one consumer there is one chain, two consumers - two chains, three consumers - six chains, four consumers - 24 chains and so on. You may recognise this sequence as factorial.

```
> cbind(n.consumers=1:n, longest.chain=sapply(cl, max), n.chains=n.chains,
factorial.n=factorial(1:n))
```

	n.consumers	longest.chain	n.chains	factorial.n
[1,]	1	1	1	1
[2,]	2	2	2	2
[3,]	3	3	6	6
[4,]	4	4	24	24
[5,]	5	5	120	120
[6,]	6	6	720	720
[7,]	7	7	5040	5040
[8,]	8	8	40320	40320

What happens if we have even more consumers?

```
> n <- 20
> cbind(n.consumers=1:n, longest.chain=1:n, factorial.n=factorial(1:n))
```

	n.consumers	longest.chain	factorial.n
[1,]	1	1	1.000000e+00
[2,]	2	2	2.000000e+00
[3,]	3	3	6.000000e+00
[4,]	4	4	2.400000e+01
[5,]	5	5	1.200000e+02
[6,]	6	6	7.200000e+02
[7,]	7	7	5.040000e+03
[8,]	8	8	4.032000e+04
[9,]	9	9	3.628800e+05
[10,]	10	10	3.628800e+06
[11,]	11	11	3.991680e+07
[12,]	12	12	4.790016e+08
[13,]	13	13	6.227021e+09
[14,]	14	14	8.717829e+10
[15,]	15	15	1.307674e+12
[16,]	16	16	2.092279e+13
[17,]	17	17	3.556874e+14

```
[18,]      18   18 6.402374e+15
[19,]      19   19 1.216451e+17
[20,]      20   20 2.432902e+18
```

The number of trophic chains quickly becomes too large to compute within practical time and within available memory, and this for communities with just a single producer. When computing chains, cheddar allocates memory as required, with a safety limit to prevent too much of the system's memory from being consumed. If you see an error message 'Unable to compute paths' then this safety limit has been reached. The limit can be altered by setting the 'cheddarMaxQueue' option.

```
> # Set to a low number to illustrate the error
> options(cheddarMaxQueue=10)
> tryCatch(TrophicChains(TL84), error=print)

<simpleError in .TrophicChainsSize(community): Unexpected error>

> # Default value
> options(cheddarMaxQueue=NULL)
> chains <- TrophicChains(TL84)
```

If you encounter this error message you can increase the value of 'cheddarMaxQueue' (from it's default of  $1e7$ ), but it is likely that the food-web is so complex that it will not be possible to compute all paths. Setting 'cheddarMaxQueue' to 0 disables the safety limit.

### 3.3.7 Trophic level

Several different measures of trophic level are used (e.g. Williams and Martinez, 2004; Jonsson et al., 2005; Zook et al., 2011.) The `PreyAveragedTrophicLevel` function uses the matrix-inversion method of Levine (1980) to compute trophic levels (Williams and Martinez, 2004). This method is very fast and accounts for flow through loops. A different measure of trophic level is offered by the `ChainAveragedTrophicLevel` function, which enumerates every unique food chain in the web (using `TrophicChainsStats`) and computes the mean position of each node in every chain (Williams and Martinez, 2004). The method of `ChainAveragedTrophicLevel` is the same as that described as 'trophic height' by Jonsson et al. (2005) and the name `TrophicHeight` is a synonym for `ChainAveragedTrophicLevel`. `ChainAveragedTrophicLevel` might be noticeably slower than `PreyAveragedTrophicLevel` for very large and/or highly connected food webs.

```
> tail(NPS(TL84, c('PreyAveragedTrophicLevel', 'ChainAveragedTrophicLevel')), 10)
```

	PreyAveragedTrophicLevel	ChainAveragedTrophicLevel
<i>Orthocyclops modestus</i>	3.205357	4.200000
<i>Ploesoma</i> sp.	2.000000	2.000000
<i>Polyarthra vulgaris</i>	2.000000	2.000000
<i>Trichocerca multicrinis</i>	2.000000	2.000000
<i>Trichocerca cylindrica</i>	2.000000	2.000000
<i>Tropocyclops prasinus</i>	3.142857	3.333333
<i>Chaoborus punctipennis</i>	3.171344	4.602527
<i>Phoxinus eos</i>	3.529951	5.168337
<i>Phoxinus neogaeus</i>	3.529951	5.168337
<i>Umbra limi</i>	3.802678	5.835003

Cheddar offers the six different measures of trophic level described by Williams and Martinez (2004). A function is provided for each one. The `TrophicLevels` convenience function returns a matrix containing all six.

```
> tail(TrophicLevels(TL84), 10)
```

	ShortestTL	ShortWeightedTL	LongestTL	LongWeightedTL
<i>Orthocyclops modestus</i>	3	3.102679	5	4.102679
<i>Ploesoma</i> sp.	2	2.000000	2	2.000000
<i>Polyarthra vulgaris</i>	2	2.000000	2	2.000000
<i>Trichocerca multicrinis</i>	2	2.000000	2	2.000000
<i>Trichocerca cylindrica</i>	2	2.000000	2	2.000000
<i>Tropocyclops prasinus</i>	3	3.071429	4	3.571429
<i>Chaoborus punctipennis</i>	2	2.585672	6	4.585672
<i>Phoxinus eos</i>	3	3.264975	7	5.264975
<i>Phoxinus neogaeus</i>	3	3.264975	7	5.264975
<i>Umbra limi</i>	3	3.401339	8	5.901339
	ChainAveragedTL	PreyAveragedTL		
<i>Orthocyclops modestus</i>	4.200000	3.205357		
<i>Ploesoma</i> sp.	2.000000	2.000000		
<i>Polyarthra vulgaris</i>	2.000000	2.000000		
<i>Trichocerca multicrinis</i>	2.000000	2.000000		
<i>Trichocerca cylindrica</i>	2.000000	2.000000		
<i>Tropocyclops prasinus</i>	3.333333	3.142857		
<i>Chaoborus punctipennis</i>	4.602527	3.171344		
<i>Phoxinus eos</i>	5.168337	3.529951		
<i>Phoxinus neogaeus</i>	5.168337	3.529951		
<i>Umbra limi</i>	5.835003	3.802678		

See the help page for `TrophicLevels` for more information on these different measures.

### 3.3.8 Link strengths

Cheddar's data format allows zero, one or many measures of link strength to be defined simply by adding additional columns to the `trophic.links.csv` file (Section 3). It is also possible to define and use theoretical measures of link strength. Cheddar's `PredationMatrix` and `FlowBasedTrophicLevel` functions allow empirical and/or theoretical link strengths to be used.

The `Benguela` dataset contains empirical estimates of diet fraction for each trophic link (Yodzis, 1998), available as the 'diet.fraction' property.

```
> head(TLPS(Benguela))
```

	resource	consumer	diet.fraction
1	Phytoplankton	Bacteria	98.8
2	Benthic filter feeders	Benthic carnivores	100.0
3	Phytoplankton	Microzooplankton	30.6
4	Bacteria	Microzooplankton	29.7
5	Microzooplankton	Microzooplankton	29.7
6	Phytoplankton	Mesozooplankton	45.0

A binary predation matrix contains just '0' and '1'.

```
> pm <- PredationMatrix(Benguela)
```

We can weight the predation matrix by empirical diet fractions.

```
> pm <- PredationMatrix(Benguela, weight='diet.fraction')
```

These matrices are 29 x 29 and so are not shown for brevity.

The `FlowBasedTrophicLevel` function uses the same matrix inversion technique as `PreyAveragedTrophicLevel` and uses the 'weight.by' node property to provide an estimate of energy flow through each trophic link. We can easily compare these two different ways of computing trophic level.

```
> cbind(PreyAveragedTrophicLevel(Benguela),  
        FlowBasedTrophicLevel(Benguela, weight.by='diet.fraction'))
```

	[,1]	[,2]
Phytoplankton	1.000000	1.000000
Benthic filter feeders	1.000000	1.000000
Bacteria	2.000000	2.000000
Benthic carnivores	2.000000	2.000000
Microzooplankton	3.000000	2.985075
Mesozooplankton	3.000000	2.992537
Macrozooplankton	3.000000	2.797015
Gelatinous zooplankton	3.250000	3.244403
Anchovy	3.500000	3.826134
Pilchard	3.333333	2.651672
Round herring	4.000000	3.914328
Lightfish	4.000000	3.875224
Lanternfish	4.000000	3.875224
Goby	3.500000	3.238903
Other pelagics	3.812500	3.952869
Horse mackerel	4.000000	3.908589
Chub mackerel	4.250000	4.165017
Other groundfish	4.776994	4.144872
Hakes	4.930366	4.767544
Squid	4.804069	4.609870
Tunas	4.844189	4.421241
Snoek	4.733434	4.713249
Kob	4.909476	4.658580
Yellowtail	4.897690	4.719394
Geelbek	4.865726	4.526048
Whales and dolphins	4.888027	4.769149
Birds	4.985516	4.737059
Seals	5.113082	4.829057
Sharks	5.183100	4.933519

Theoretical per capita interaction strengths can be computed from body-mass ratios raised to a power, typically taken to be 2/3 or 3/4 (Emmerson and Raffaelli, 2004; Reuman and Cohen, 2005; Layer et al., 2010). We can define a function to compute these theoretical interaction strengths and use it in computation of trophic levels.

```

> InteractionStrength <- function(community)
{
  tlps <- TLPS(community, node.properties='M')
  return ((tlps$consumer.M / tlps$resource.M)^3/4)
}
> # The InteractionStrength() function can be used together with TLPS() to
> # compute the theoretical interaction strength between each resource-consumer pair
> head(TLPS(Benguela, link.properties='InteractionStrength'))

```

	resource	consumer	InteractionStrength
1	Phytoplankton	Bacteria	2.5e-13
2	Benthic filter feeders	Benthic carnivores	2.5e-01
3	Phytoplankton	Microzooplankton	2.5e-01
4	Bacteria	Microzooplankton	2.5e+11
5	Microzooplankton	Microzooplankton	2.5e-01
6	Phytoplankton	Mesozooplankton	2.5e+05

We can use this measure of interaction strength to compute flow-based trophic level.

```

> cbind(PreyAveragedTrophicLevel(Benguela),
         FlowBasedTrophicLevel(Benguela, weight.by='diet.fraction'),
         FlowBasedTrophicLevel(Benguela, weight.by='InteractionStrength'))

```

	[,1]	[,2]	[,3]
Phytoplankton	1.000000	1.000000	1.000000
Benthic filter feeders	1.000000	1.000000	1.000000
Bacteria	2.000000	2.000000	2.000000
Benthic carnivores	2.000000	2.000000	2.000000
Microzooplankton	3.000000	2.985075	3.000000
Mesozooplankton	3.000000	2.992537	3.000000
Macrozooplankton	3.000000	2.797015	2.000002
Gelatinous zooplankton	3.250000	3.244403	3.000000
Anchovy	3.500000	3.826134	3.000000
Pilchard	3.333333	2.651672	2.000002
Round herring	4.000000	3.914328	3.999999
Lightfish	4.000000	3.875224	3.999999
Lanternfish	4.000000	3.875224	3.999999
Goby	3.500000	3.238903	3.000000
Other pelagics	3.812500	3.952869	3.999999
Horse mackerel	4.000000	3.908589	3.999999
Chub mackerel	4.250000	4.165017	3.999999
Other groundfish	4.776994	4.144872	3.999999
Hakes	4.930366	4.767544	3.999999
Squid	4.804069	4.609870	3.018712
Tunas	4.844189	4.421241	4.796740
Snoek	4.733434	4.713249	3.024660
Kob	4.909476	4.658580	3.000829
Yellowtail	4.897690	4.719394	3.000830
Geelbek	4.865726	4.526048	3.453351

Whales and dolphins	4.888027	4.769149	3.006892
Birds	4.985516	4.737059	3.999999
Seals	5.113082	4.829057	4.796740
Sharks	5.183100	4.933519	3.999999

The weighting functionality offered by the `PredationMatrix` and `FlowBasedTrophicLevel` functions make it possible to use multiple empirical and/or theoretical link strengths and interaction strengths in analyses.

## 4 Community manipulations

### 4.1 Node order

The ordering of nodes within a community can be important both for presentation and analysis. Cheddar's `OrderCommunity` function reorders nodes and returns a new community object. `OrderCommunity` accepts names that meets the criteria of the `properties` parameter of the `NPS` function. This includes the names of 'first-class' properties, such as `M`, and the names of functions that take a single community and return a value for each node, such as `Degree`, which returns the number of trophic links for each node. The following examples order TL84 by increasing body mass and by increasing degree.

```
> TL84.increasing.M <- OrderCommunity(TL84, 'M', title='Increasing M')
> head(NPS(TL84.increasing.M, c('M', 'Degree')))
```

	M	Degree
Chromulina sp.	3.03e-14	18
Dactylococcopsis fascicularis	1.32e-13	4
Diceras sp.	1.53e-13	0
Trachelomonas sp.	1.75e-13	18
Cryptomonas sp. 1	2.03e-13	18
Closteriopsis longissimus	2.37e-13	3

```
> TL84.increasing.degree <- OrderCommunity(TL84, 'Degree',
                                               title='Increasing degree')
> head(NPS(TL84.increasing.degree, c('M', 'Degree')))
```

	M	Degree
Asterionella formosa	1.12e-12	0
Chrysosphaerella longispina	8.31e-10	0
Diceras sp.	1.53e-13	0
Rhizosolenia sp.	6.86e-13	0
Spinocosmarium sp.	3.71e-12	0
Staurastrum sp.	4.30e-12	0

Similar to R's `order` function, `OrderCommunity` can sort by more than one name with subsequent names used to break ties. We can use this to sort alphabetically by category and then by increasing `M` within each category.

```
> TL84.category.then.M <- OrderCommunity(TL84, 'category', 'M')
> head(NPS(TL84.category.then.M, c('category', 'M')))
```

	category	M
Keratella cochlearis	invertebrate	1.00e-11
Keratella testudo	invertebrate	1.00e-11
Kellicottia sp.	invertebrate	2.00e-11
Conochilus (solitary)	invertebrate	3.50e-11
Ploesoma sp.	invertebrate	1.05e-10
Gastropus stylifer	invertebrate	1.35e-10

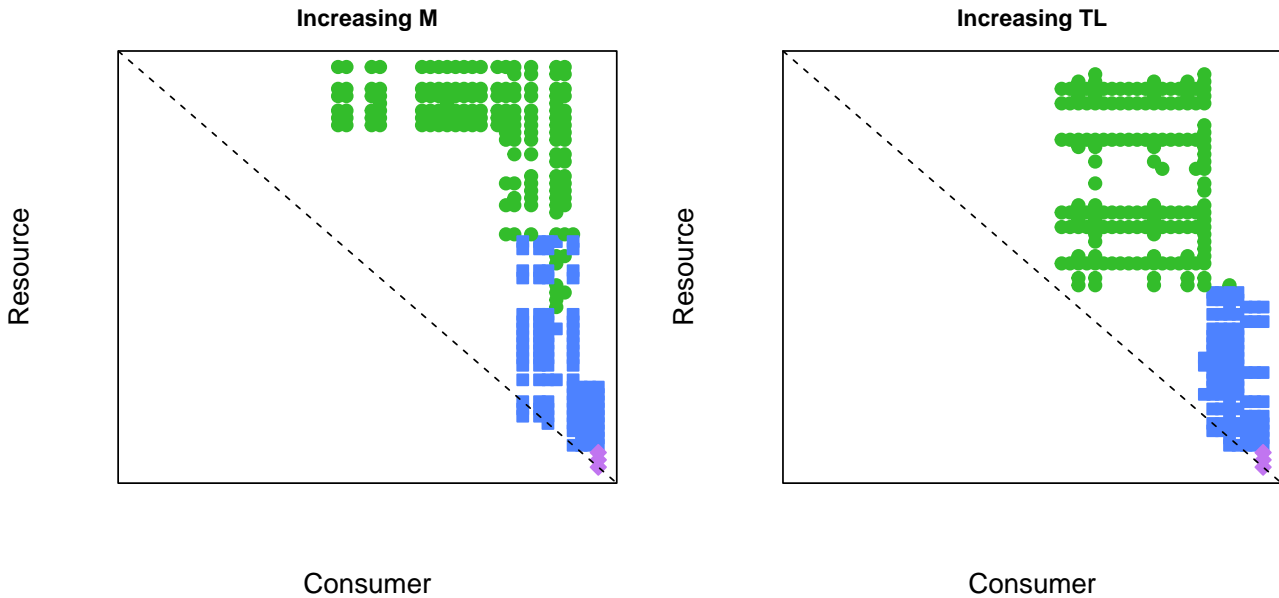
## 4.2 Node order and intervality

Visualising the food web as a predation matrix is central to many analyses and theories. There has been much recent interest in the relationship between food web structure and species' niches, in particular the role of body size on determining a species' position in a food web and the effect on intervality - a measure of the adjacency of resources and consumers in the food web (Williams and Martinez, 2000; Stouffer et al., 2006; Zook et al., 2011). We can use `OrderCommunity` to explore the effect ordering species along different niche axes. The code fragment below creates two new orderings of TL84, one by increasing body mass and the other by increasing trophic level, with random ordering within ties for trophic level (Zook et al., 2011).

```
> # Increasing M
> TL84.increasing.M <- OrderCommunity(TL84, 'M', title='Increasing M')
> new.order <- order(PreyAveragedTrophicLevel(TL84), sample(1:56))
> TL84.increasing.TL <- OrderCommunity(TL84, new.order=new.order,
                                         title='Increasing TL')
```

We could use any of Cheddar's different measure of trophic level (Section 3.3.7). The `PlotPredationMatrix` function allows us to graphically compare the effect of these different orderings.

```
> par(mfrow=c(1, 2))
> PlotPredationMatrix(TL84.increasing.M)
> PlotPredationMatrix(TL84.increasing.TL)
```



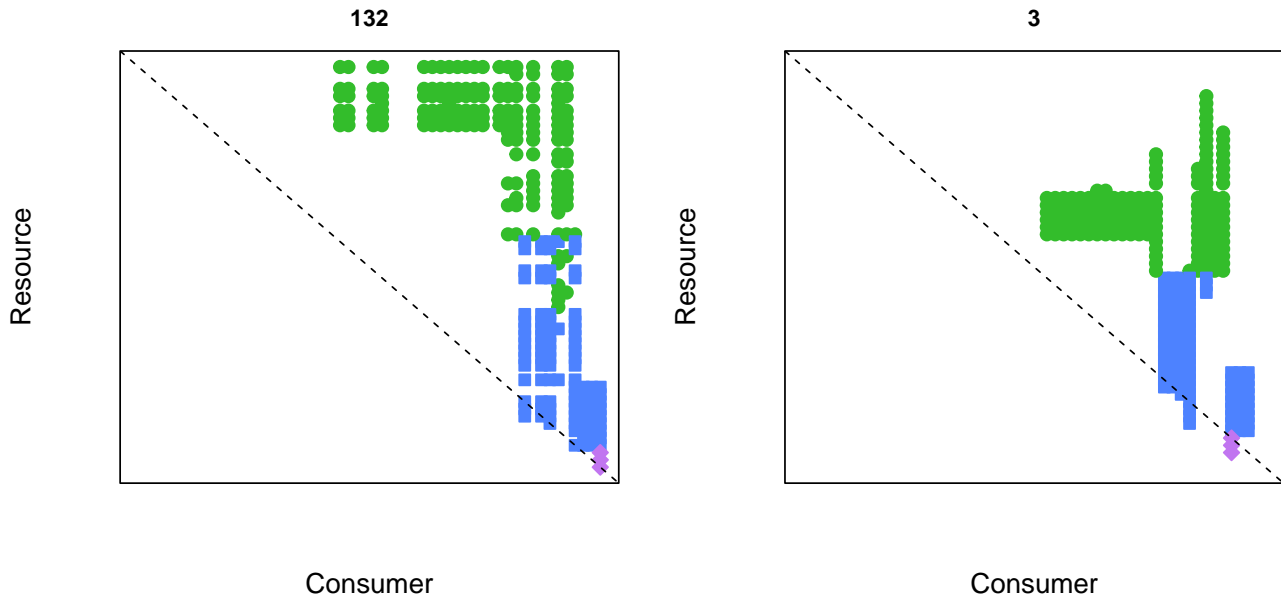
The total number of gaps in diets (columns) and consumers (rows) (Stouffer et al., 2011; Zook et al., 2011):

```
> SumDietGaps(TL84.increasing.M)
[1] 132
> SumDietGaps(TL84.increasing.TL)
```

```
[1] 366
> SumConsumerGaps(TL84.increasing.M)
[1] 154
> SumConsumerGaps(TL84.increasing.TL)
[1] 158
```

The `MinimiseSumDietGaps` function implements simulated annealing learning to minimise `SumDietGaps`, as described by Stouffer et al. (2006). Simulated annealing learning is a stochastic method so several optimisations might be required to find the global minimum; this is done by setting the ‘n’ parameter greater than 1:

```
> par(mfrow=c(1, 2))
> PlotPredationMatrix(TL84.increasing.M, main=SumDietGaps(TL84.increasing.M))
> res <- MinimiseSumDietGaps(TL84, n=10)
> PlotPredationMatrix(res$reordered, main=SumDietGaps(res$reordered))
```



`MinimiseSumConsumerGaps` uses the same method to minimise the gaps in each species' consumers (Zook et al., 2011).

```
> par(mfrow=c(1, 2))
> PlotPredationMatrix(TL84.increasing.M,
  main=paste('Ordered by M - sum consumer gaps',
  SumConsumerGaps(TL84.increasing.M)))
> res <- MinimiseSumConsumerGaps(TL84, n=10)
> PlotPredationMatrix(res$reordered,
  main=paste('Optimised - sum consumer gaps',
  SumConsumerGaps(res$reordered)))
```



### 4.3 Removing nodes

Isolated nodes are often removed from food-web analyses (e.g. Jonsson et al., 2005). `RemoveIsolatedNodes` is a convenience function that returns a new `Community` with isolated nodes removed.

```
> NumberOfNodes(TL84)
[1] 56

> IsolatedNodes(TL84)
[1] "Asterionella formosa"      "Chrysosphaerella longispina"
[3] "Diceras sp."              "Rhizosolenia sp."
[5] "Spinocosmarium sp."       "Staurastrum sp."

> NumberOfTrophicLinks(TL84)
[1] 269
```

```

> TL84.no.isolated <- RemoveIsolatedNodes(TL84)
> NumberOfNodes(TL84.no.isolated)           # Six fewer species
[1] 50

> IsolatedNodes(TL84.no.isolated)          # No isolated species
character(0)

> NumberOfTrophicLinks(TL84.no.isolated)    # Number of trophic links unchanged
[1] 269

The general-purpose RemoveNodes function returns a new Community object with one or more nodes removed.

> NumberOfNodes(TL84)
[1] 56

> NumberOfTrophicLinks(TL84)
[1] 269

> # Remove the first ten nodes
> TL84.r <- RemoveNodes(TL84, 1:10)
> NumberOfNodes(TL84.r)
[1] 46

> NumberOfTrophicLinks(TL84.r)
[1] 213

> # Remove producers
> TL84.r <- RemoveNodes(TL84, 'producer'==NP(TL84, 'category'))
> NumberOfNodes(TL84.r)
[1] 25

> NumberOfTrophicLinks(TL84.r)
[1] 103

> # Remove species by name
> to.remove <- c("Cryptomonas sp. 1", "Chroococcus dispersus",
   "Unclassified flagellates", "Chromulina sp.",
   "Selenastrum minutum", "Trachelomonas sp.")
> TL84.r <- RemoveNodes(TL84, to.remove)
> NumberOfNodes(TL84.r)
[1] 50

```

```

> NumberOfTrophicLinks(TL84.r)
[1] 161

> # Three different ways of removing node 56 (Umbra limi)
> TL84.ra <- RemoveNodes(TL84, 56)
> TL84.rb <- RemoveNodes(TL84, 'Umbra limi')
> TL84.rc <- RemoveNodes(TL84, c(rep(FALSE,55), TRUE))
> identical(TL84.ra, TL84.rb) # TRUE

[1] TRUE

> identical(TL84.ra, TL84.rc) # TRUE

[1] TRUE

```

The `RemoveNodes` function takes an argument called ‘method’, which indicates how removals should be propagated through the food web. If ‘method’ is ‘direct’ (the default), only the nodes in `remove` are removed. If ‘method’ is ‘secondary’, secondarily extinct nodes - those that directly consume one or more nodes in ‘remove’ and that no longer have any resources (except themselves) after the removal - are also removed. If ‘method’ is ‘cascade’, a multistep version of ‘secondary’ is applied. This has the effect of propagating extinctions through the community - all consumers that are ultimately dependent upon all species in ‘remove’, and upon no other nodes (except themselves), will be removed.

```

> # The behaviours of the different methods
> NumberOfNodes(TL84)           # 56 nodes in total

[1] 56

> length(BasalNodes(TL84))     # 25 basal nodes

[1] 25

> length(IsolatedNodes(TL84)) # 6 isolated nodes

[1] 6

> RemoveNodes(TL84, BasalNodes(TL84)) # 56 - 25 = 31 nodes remain

```

Tuesday Lake sampled in 1984 (25 nodes directly removed) containing 31 nodes and 103 trophic links

```
> RemoveNodes(TL84, BasalNodes(TL84), method='secondary') # 14 nodes remain
```

Tuesday Lake sampled in 1984 (25 nodes directly removed) containing 14 nodes and 30 trophic links

```
> RemoveNodes(TL84, BasalNodes(TL84), method='cascade')    # The 6 isolated nodes remain
```

Tuesday Lake sampled in 1984 (25 nodes directly removed) containing 6 nodes

#### 4.4 Removing cannibalistic links

`RemoveCannibalisticLinks` returns a new `Community` without those trophic links in which a node consumes itself.

```
> NumberOfNodes(TL84)

[1] 56

> Cannibals(TL84)          # 5 species

[1] "Cyclops varians rubellus" "Orthocyclops modestus"    "Tropocyclops prasinus"
[4] "Chaoborus punctipennis"   "Umbra limi"

> NumberOfTrophicLinks(TL84)

[1] 269

> TL84.no.cannibals <- RemoveCannibalisticLinks(TL84)
> NumberOfNodes(TL84.no.cannibals)           # Number of nodes unchanged

[1] 56

> Cannibals(TL84.no.cannibals)            # No species

character(0)

> NumberOfTrophicLinks(TL84.no.cannibals)  # 5 fewer trophic links

[1] 264
```

#### 4.5 Lumping nodes

Certain analyses call for food-web nodes to be merged. In order to reduce biases, nodes that share the same resources and consumers, so-called ‘trophic species’, are commonly lumped together (Briand and Cohen, 1984; Pimm et al., 1991; Williams and Martinez, 2000). The `LumpTrophicSpecies` performs this task and returns a new `Community` object.

```
> NumberOfNodes(TL84)

[1] 56

> TL84.lumped <- LumpTrophicSpecies(TL84)
> length(unique(TrophicSpecies(TL84)))      # 22 trophic species in TL84...

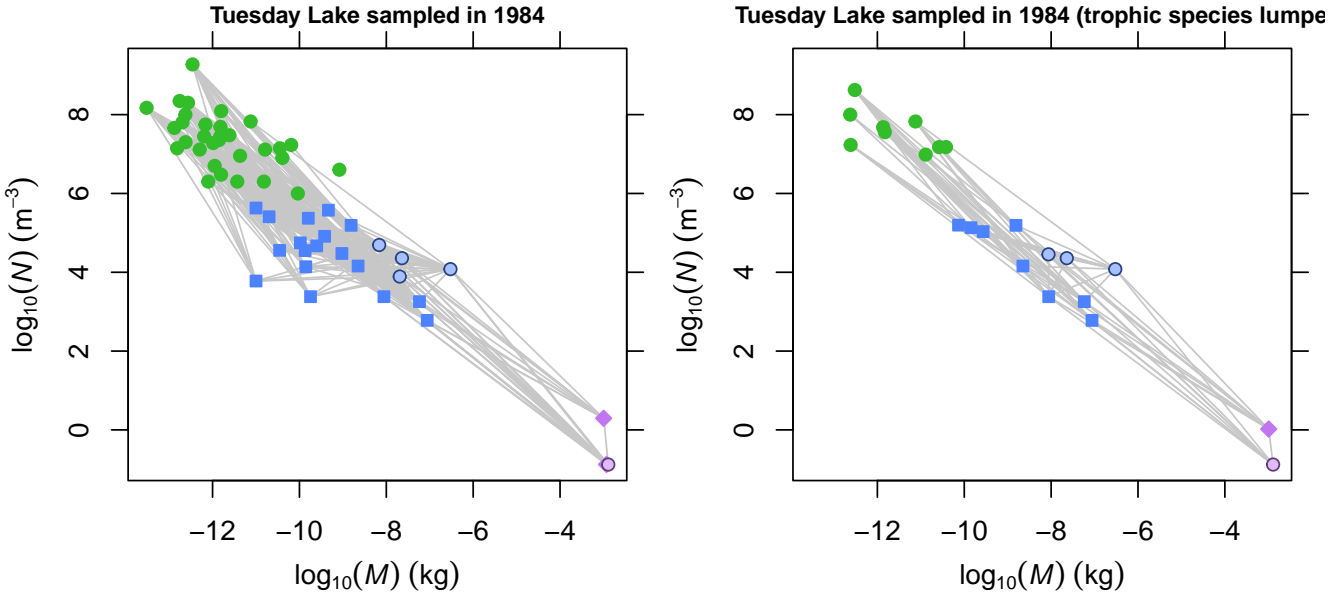
[1] 22

> NumberOfNodes(TL84.lumped)                  # ... and 22 nodes in the lumped web

[1] 22
```

The plot below shows the lumped and unlumped webs.

```
> par(mfrow=c(1, 2))
> plot(TL84)
> plot(TL84.lumped, xlim=range(Log10M(TL84)), ylim=range(Log10N(TL84)))
```



The `LumpNodes` function is a more general-purpose function that allows any nodes in a community to be lumped together. It takes a parameter ‘lump’, which should be a vector of length `NumberOfNodes`. Nodes with the same value of ‘lump’ will be merged together. This example lumps together isolated species in TL84.

```
> length(which(IsIsolatedNode(TL84))) # 6 isolated species
[1] 6

> IsolatedNodes(TL84) # Names of isolated nodes
[1] "Asterionella formosa"      "Chrysosphaerella longispina"
[3] "Diceras sp."              "Rhizosolenia sp."
[5] "Spinocosmarium sp."       "Staurastrum sp."

> lump <- NP(TL84, 'node')      # Existing node names
> # Give isolated nodes the same lump value
> lump[IsolatedNodes(TL84)] <- 'Isolated nodes lumped together'
> TL84.lumped <- LumpNodes(TL84, lump)
> NumberOfNodes(TL84)          # 56 nodes in unlumped web
[1] 56

> NumberOfNodes(TL84.lumped) # 51 nodes in lumped web
```

```
[1] 51

> IsolatedNodes(TL84.lumped) # A single node
[1] "Isolated nodes lumped together"
```

By default, numeric values are weighted by numerical abundance,  $N$ .

This trivial example shows that no nodes are lumped if values in lump are unique to each node.

```
> lump <- NP(TL84, 'node')
> identical(TL84, LumpNodes(TL84, lump, title=CP(TL84, 'title')))

[1] FALSE
```

The Ythan Estuary dataset contains two species that are split by life stage: *Platichthys flesus* (european flounder) and *Somateria mollissima* (common eider). The code fragment below shows how to lump these in to a single node for each species.

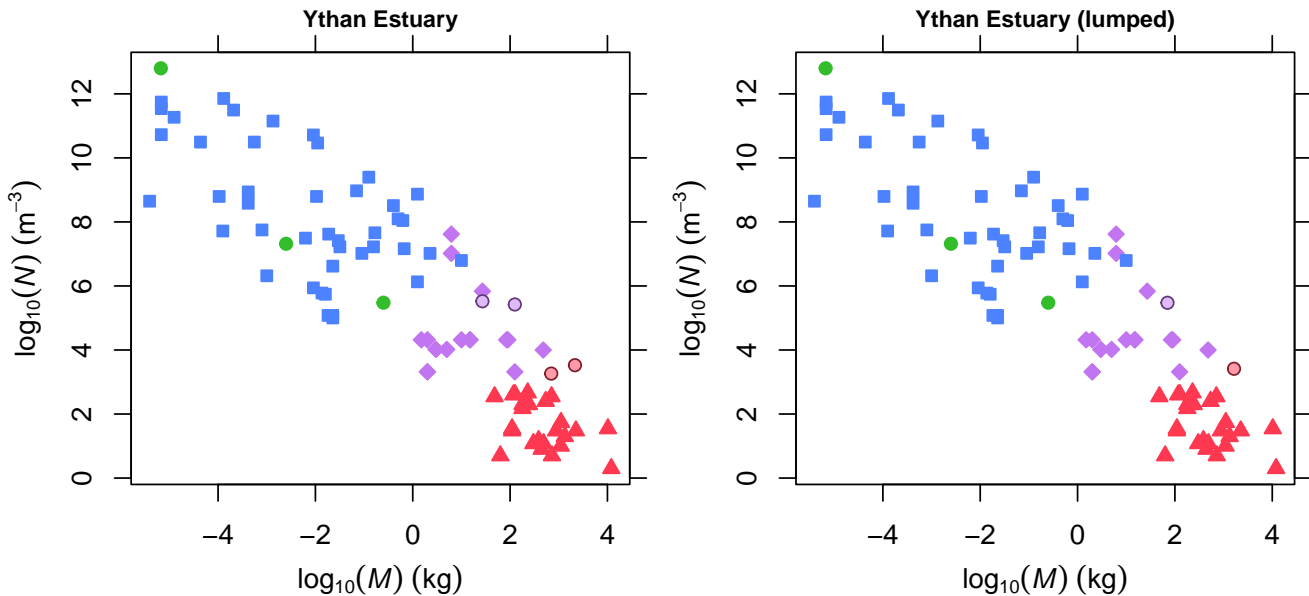
```
> data(YthanEstuary)
> # The names of nodes in YthanEstuary
> lump <- NP(YthanEstuary, 'node')
> # European flounder:
> # "Platichthys flesus" and "Platichthys flesus (juvenile)"
> # Lump these in to one node
> lump["Platichthys flesus (juvenile)"==lump] <- "Platichthys flesus"
> # Common eider:
> # "Somateria mollissima" and "Somateria mollissima (juvenile)"
> # Lump these in to one node
> lump["Somateria mollissima (juvenile)"==lump] <- "Somateria mollissima"
> YthanEstuary.lumped <- LumpNodes(YthanEstuary, lump)
> NumberOfNodes(YthanEstuary)      # 92

[1] 92

> NumberOfNodes(YthanEstuary.lumped) # 90
[1] 90
```

Graphically compare the two communities.

```
> # Plot the original and lumped communities
> par(mfrow=c(1,2))
> plot(YthanEstuary, highlight.nodes=c("Platichthys flesus",
+ "Platichthys flesus (juvenile)",
+ "Somateria mollissima",
+ "Somateria mollissima (juvenile)",
+ show.web=FALSE)
> plot(YthanEstuary.lumped, highlight.nodes=c("Platichthys flesus",
+ "Somateria mollissima"),
+ show.web=FALSE)
```



The default behaviour of `LumpNodes` and `LumpTrophicSpecies` is to aggregate numeric node properties by computing the  $N$ -weighted mean.

```
> NPS(YthanEstuary.lumped)[["Platichthys flesus", c('M', 'N')]]
```

	M	N
Platichthys flesus	70.36585	298480

```
> # These values were computed as follows
> nps <- NPS(YthanEstuary)
> M <- nps[c("Platichthys flesus", "Platichthys flesus (juvenile)", 'M')]
> N <- nps[c("Platichthys flesus", "Platichthys flesus (juvenile)", 'N')]
> # Arithmetic mean of N
> mean(N)
[1] 298480
```

```

> # N-weighted mean of M
> weighted.mean(M, N)

[1] 70.36585

```

The ‘weight.by’ parameter controls this behaviour:

```

> YthanEstuary.lumped2 <- LumpNodes(YthanEstuary, lump, weight.by=NULL)
> NPS(YthanEstuary.lumped2)[["Platichthys flesus", c('M', 'N')]

          M         N
Platichthys flesus 76 298480

```

```

> # Computed as the arithmetic means of M and N
> mean(M)

```

```
[1] 76
```

```
> mean(N)
```

```
[1] 298480
```

```
>
```

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