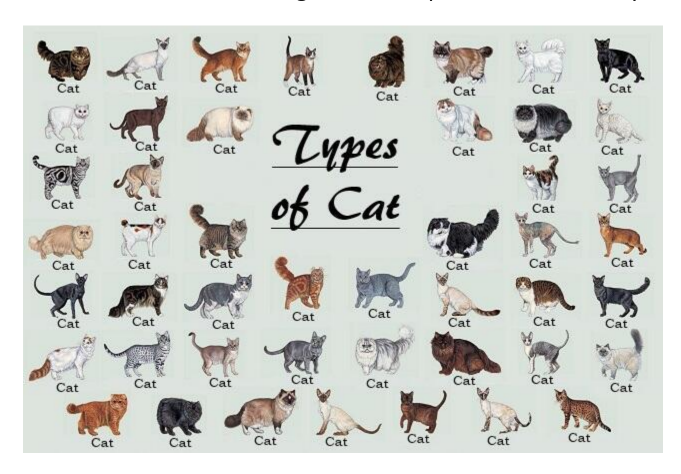
Characterizing insect communities using measures of diversity

ENTMLGY 6702 Entomological Techniques and Data Analysis



Learning objectives

- 1) Distinguish among different diversity metrics
- 2) Understand how multiple dimensions of diversity contribute to characterizing communities
- 3) Interpret the outputs of different metrics

Traditional approach has focused on taxonomy

Measure unique "types" and compare among samples or communities

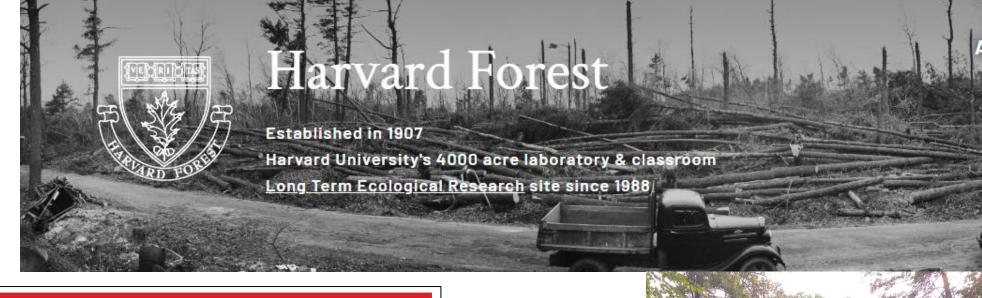
<u>Type</u>: taxonomic unit (species, genera, families, orders)

- 1. Abundance measures the number of individuals of each distinct type
- 2. Richness measures the number of distinct types
- 3. Evenness measures the relative abundance of each type
- 4. Diversity measures the number of types and their evenness

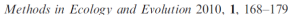
- Hemlock girdled
- Hemlock logged
- Hemlock control
- Hardwood control

n = 2

90 x 90 m plots



Methods in Ecology and Evolution



doi: 10.1111/j.2041-210X.2010.00025.x

Experimentally testing the role of foundation species in forests: the Harvard Forest Hemlock Removal **Experiment**

Aaron M. Ellison*, Audrey A. Barker-Plotkin, David R. Foster and David A. Orwig

Harvard Forest, Harvard University, 324 North Main Street, Petersham, MA 01366, USA





Long format

~		J										
•	year [‡]	date [‡]	block [‡]	plot [‡]	treatment [‡]	moose.cage [‡]	trap.type [‡]	trap.num [‡]	genus [‡]	species	code [‡]	abundance [‡]
1	2003	2003-06-01	Ridge	4	Logged	NA	bait	1 hour	Aphaenogaster	fulva	aphful	2
2	2003	2003-06-01	Ridge	4	Logged	NA	bait	1 hour	Aphaenogaster	picea	aphpic	2
3	2003	2003-06-01	Ridge	4	Logged	NA	bait	1 hour	Temnothorax	longispinosus	temion	1
4	2003	2003-06-01	Ridge	4	Logged	NA	bait	15 minutes	Aphaenogaster	picea	aphpic	2
5	2003	2003-06-01	Ridge	4	Logged	NA	hand	NA	Aphaenogaster	picea	aphpic	1
6	2003	2003-06-01	Ridge	4	Logged	NA	hand	NA	Camponotus	pennsylvanicus	campen	1
7	2003	2003-06-01	Ridge	4	Logged	NA	hand	NA	Lasius	americanus	lasame	1
8	2003	2003-06-01	Ridge	4	Logged	NA	hand	NA	Myrmica	punctiventris	myrpun	1
9	2003	2003-06-01	Ridge	4	Logged	NA	hand	NA	Temnothorax	longispinosus	temion	1
10	2003	2003-06-01	Ridge	4	Logged	NA	litter	1	Aphaenogaster	fulva	aphful	1
11	2003	2003-06-01	Ridge	4	Logged	NA	litter	1	Aphaenogaster	picea	aphpic	1
12	2003	2003-06-01	Ridge	4	Logged	NA	litter	2	Aphaenogaster	picea	aphpic	1
13	2003	2003-06-01	Ridge	4	Logged	NA	litter	2	Temnothorax	longispinosus	temion	2
14	2003	2003-06-01	Ridge	4	Logged	NA	litter	3	Aphaenogaster	picea	aphpic	1
15	2003	2003-06-01	Ridge	4	Logged	NA	pitfall	1	Aphaenogaster	picea	aphpic	1
16	2003	2003-06-01	Ridge	4	Logged	NA	pitfall	15	Aphaenogaster	picea	aphpic	1
17	2003	2003-06-01	Ridge	4	Logged	NA	pitfall	17	Aphaenogaster	picea	aphpic	2
18	2003	2003-06-01	Ridge	4	Logged	NA	pitfall	19	Aphaenogaster	picea	aphpic	1
19	2003	2003-06-01	Ridge	4	Logged	NA	pitfall	2	Aphaenogaster	picea	aphpic	1
20	2003	2003-06-01	Ridge	4	Logged	NA	pitfall	20	Temnothorax	longispinosus	temion	1
21	2003	2003-06-01	Ridge	4	Logged	NA	pitfall	22	Aphaenogaster	picea	aphpic	1
22	2003	2003-06-01	Ridge	4	Logged	NA	pitfall	23	Aphaenogaster	picea	aphpic	1
23	2003	2003-06-01	Ridge	4	Logged	NA	pitfall	24	Aphaenogaster	picea	aphpic	1
24	2003	2003-06-01	Ridge	4	Logged	NA	pitfall	5	Aphaenogaster	picea	aphpic	2
25	2003	2003-06-01	Ridge	4	Logged	NA	pitfall	7	Aphaenogaster	picea	aphpic	1
26	2003	2003-06-01	Ridge	5	Girdled	NA	bait	1 hour	Aphaenogaster	fulva	aphful	1

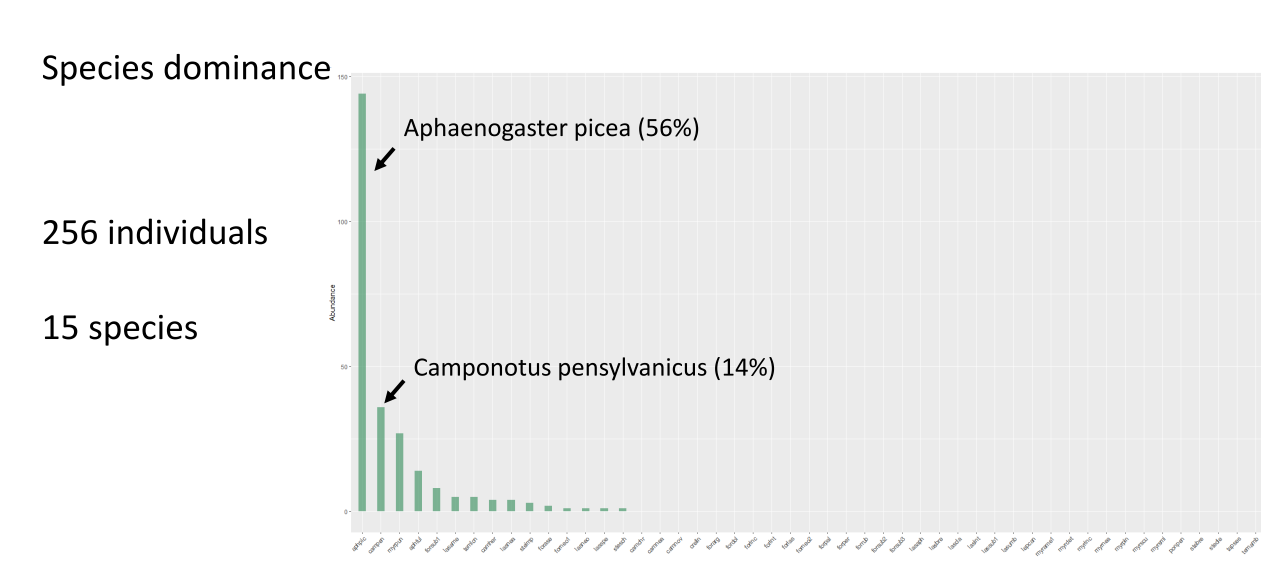
Wide format

2006

^	year [‡]	block [‡]	plot [‡]	treatment [‡]	trap.type [‡]	aphful [‡]	aphpic [‡]	camchr [‡]	camher [‡]	camnea [‡]	camnov [‡]	campen [‡]	crelin [‡]	forarg [‡]	forase	fordo
1	2006	Ridge	4	Logged	bait	0	2	0	0	0	0	2	0	0	0	
2	2006	Ridge	4	Logged	hand	0	2	0	2	0	0	2	0	0	0	
3	2006	Ridge	4	Logged	litter	0	1	0	0	0	0	0	0	0	0	
4	2006	Ridge	4	Logged	pitfall	0	11	0	0	0	0	1	0	0	0	
5	2006	Ridge	5	Girdled	bait	2	2	0	0	0	0	2	0	0	0	
6	2006	Ridge	5	Girdled	hand	0	1	0	0	0	0	1	0	0	0	
7	2006	Ridge	5	Girdled	litter	0	1	0	0	0	0	0	0	0	0	
8	2006	Ridge	5	Girdled	pitfall	4	19	0	0	0	0	13	0	0	0	
9	2006	Ridge	6	HemlockControl	bait	0	2	0	0	0	0	0	0	0	0	
10	2006	Ridge	6	HemlockControl	hand	0	2	0	0	0	0	0	0	0	0	
11	2006	Ridge	6	HemlockControl	litter	0	1	0	0	0	0	0	0	0	0	
12	2006	Ridge	6	HemlockControl	pitfall	1	22	0	0	0	0	0	0	0	0	
13	2006	Ridge	7	HardwoodControl	bait	0	2	0	0	0	0	2	0	0	0	
14	2006	Ridge	7	HardwoodControl	hand	0	1	0	0	0	0	1	0	0	0	
15	2006	Ridge	7	HardwoodControl	litter	0	1	0	0	0	0	0	0	0	0	
16	2006	Ridge	7	HardwoodControl	pitfall	0	18	0	0	0	0	2	0	0	0	(
17	2006	Valley	1	Girdled	bait	1	2	0	0	0	0	1	0	0	0	
18	2006	Valley	1	Girdled	hand	0	2	0	0	0	0	1	0	0	0	(
19	2006	Valley	1	Girdled	litter	0	1	0	0	0	0	0	0	0	0	
20	2006	Valley	1	Girdled	pitfall	1	10	0	0	0	0	2	0	0	0	(
21	2006	Valley	2	Logged	bait	1	2	0	0	0	0	1	0	0	0	(
22	2006	Valley	2	Logged	hand	0	1	0	2	0	0	2	0	0	0	
23	2006	Valley	2	Logged	pitfall	0	14	0	0	0	0	0	0	0	0	
24	2006	Valley	3	HemlockControl	bait	0	1	0	0	0	0	0	0	0	0	
25	2006	Valley	3	HemlockControl	hand	1	0	0	0	0	0	1	0	0	0	
26	2006	Valley	3	HemlockControl	pitfall	0	2	0	0	0	0	0	0	0	0	
27	2006	Valley	8	HardwoodControl	bait	0	1	0	0	0	0	1	0	0	0	
28	2006	Valley	8	HardwoodControl	hand	1	2	0	0	0	0	0	0	0	0	
29	2006	Valley	8	HardwoodControl	litter	0	0	0	0	0	0	0	0	0	0	

Abundance: the number of individuals of each distinct type

Descriptive information about a community is valuable



Abundance: the number of individuals of each distinct type

Indicator species analyses

Relationship between species abundance and environmental classification of sites

- 1. Reflect the biotic or abiotic state of the environment
- 2. Provide evidence for the impacts of environmental change
- 3. Predict the diversity of other species, taxa, or communities within an area

Can be used for conservation monitoring purposes

Abundance: the number of individuals of each distinct type

> indval <- multipatt(ants3[,6:20], ants2[,4], duleg = TRUE, control = how(nperm=999))</pre>

Indicator species analyses

```
> summary(indval, indvalcomp = TRUE)
Multilevel pattern analysis
 Association function: IndVal.q
 Significance level (alpha): 0.05
Total number of species: 15
 Selected number of species: 3
 Number of species associated to 1 group: 3
 Number of species associated to 2 groups: 0
Number of species associated to 3 groups: 0
List of species associated to each combination:
Group HardwoodControl #sps. 3
                   B stat p.value
                                         Myrmica punctiventris
myrpun 0.9211 1.0000 0.960 0.001 ***
                                         Formica subaenescens
forsub1 0.8596 0.5000 0.656 0.023 *
                                         Lasius americanus
lasame 0.7778 0.5000 0.624 0.029 *
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' '1
```

Indicator value index:

A = <u>specificity</u>; given that the species is found, probability that the site belongs to the target group

B = <u>sensitivity</u>; probability of finding the species in sites belonging to the site group

R package indicspecies

Species richness

How many species (or genera, families, etc)?

Absolute species richness: Count all species. Possible with some taxa, like large mammals. Very challenging with insects!

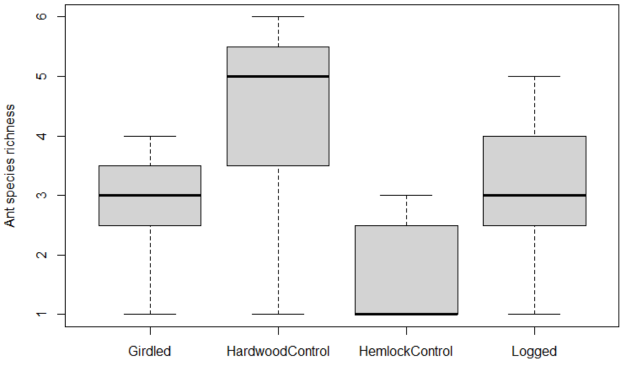
Estimate species richness by collecting samples

- 1. Numerical richness: number of species per specified number of individuals, biomass, or collection traps that do not yield density measure (such as pan traps, sticky traps, pitfall traps)
- 2. Density based richness: number of species per area or unit (Number of species per meter square, species per sweep or per flower, etc.)

Richness: the number of distinct types

Count the number of species collected per site, trap, etc.





Richness: the number of distinct types

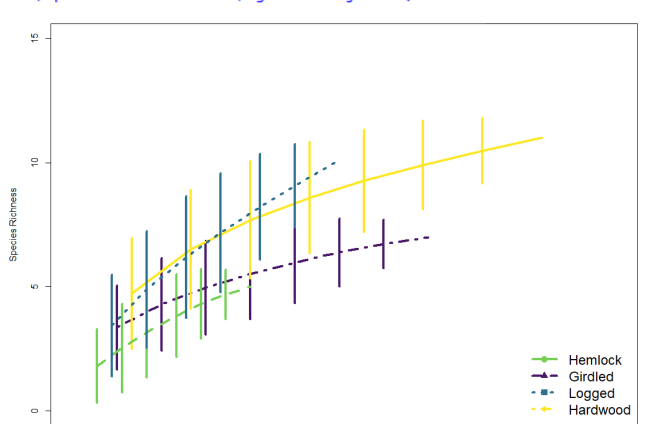
Estimate species richness with accumulation curves

```
> sp.girdled <- specaccum(girdled[6:20], method = "rarefaction", permutations = 100, gamma = "jack2")
> sp.hardwood <- specaccum(hardwood[6:20], method = "rarefaction", permutations = 100, gamma = "jack2")
> sp.hemlock <- specaccum(hemlock[6:20], method = "rarefaction", permutations = 100, gamma = "jack2")
> sp.logged <- specaccum(logged[6:20], method = "rarefaction", permutations = 100, gamma = "jack2")
```

Estimate species richness by standardizing based on:

- 1) Sites
- 2) Individuals

Permutation procedure that subsamples the data



R package vegan

Richness: the number of distinct types

Calculate first- and second-order jackknife estimates of species richness

lackknife (1st order)	$S_{p} = S_{o} + a_{1} (N-1)/N$
lackknife (2nd order)	$S_p = S_0 + a_1 (2N - 3)/N - a_2 (N - 2)^2/N(N - 1)$

Treatment	Observed species	First-order jackknife	First-order	Second-order jackknife	Second-order
	richness	estimate	%	estimate	%
Hemlock	5	6.9	72.4	6.8	73.5
Girdled	7	8.9	78.6	8.9	78.6
Logged	10	15.8	63.2	20.3	49.2
Hardwood	11	14.9	73.8	17.7	62.1



Diversity: the number of types and their evenness

Many metrics that are used to calculate diversity

- 1) Shannon index
- 2) Simpson index
- 3) Effective number of species
- 4) Hill numbers

1. Shannon – Wiener Index (also called Shannon Index, Shannon Weaver)

$$H' = -\sum_{i=1}^{n} p_i \ln p_i$$

 $p_i = n_i / N$, where n_i is the abundance of the ith species and N is the total abundance.

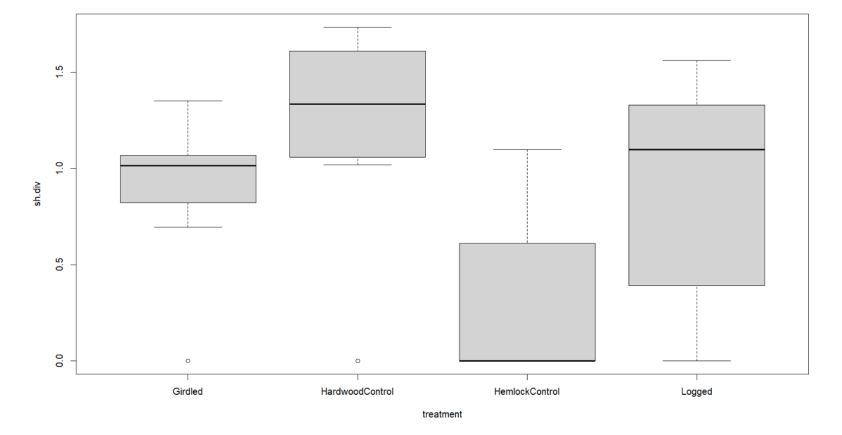
H' increases as diversity increases

You can compare H' values among sites you sample using the same method and equal effort.

1. Shannon – Wiener Index

```
> # Shannon index
> diversity(ants3[,6:20], index = "shannon")
[1] 1.3296613 1.0986123 0.0000000 0.5359610 1.3517840 1.0986123 0.0000000 0.9492451 0.0000000
[10] 1.0397208 0.0000000 0.1788449 1.3517840 1.6094379 1.0986123 1.0195341 1.0397208 1.0397208
[19] 0.6931472 0.9881542 1.5607104 1.3296613 0.2449300 0.0000000 1.0986123 0.0000000 1.6094379
[28] 1.7328680 0.0000000 1.3189601
```





2. Simpsons Diversity Index

Simpson (1949) gave the probability of any two individuals drawn at random from an infinitely large community belonging to the same species as:

$$D = \sum \frac{n_i(n_i - 1)}{N(N - 1)}$$

Gini-Simpson index

$$GS = 1-D$$

Where ni= the number of individuals in the ith species and N=the total number of individuals.

Note: *D* represents the probability of two individuals belonging to the same species, so diversity actually decreases as *D* increases.

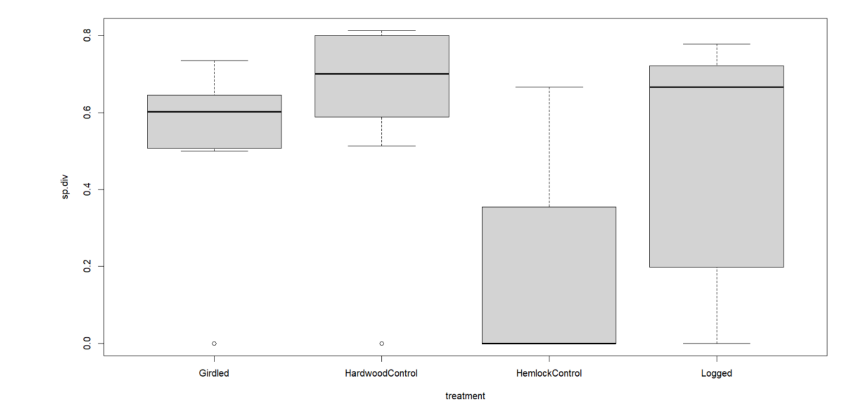
2. Simpsons Diversity Index

```
Gini-Simpson index
```

$$GS = 1-D$$

```
R package
vegan
```

```
> # Simpson index
> diversity(ants3[,6:20], index = "simpson")
[1] 0.7222222 0.6666667 0.0000000 0.2721893 0.7346939 0.6666667 0.0000000 0.5787037 0.0000000
[10] 0.6250000 0.0000000 0.0831758 0.7346939 0.8000000 0.6666667 0.5130316 0.6250000 0.5250000
[19] 0.5000000 0.5155556 0.7777778 0.7222222 0.1244444 0.0000000 0.6666667 0.0000000 0.8000000
[28] 0.8125000 0.0000000 0.6640368
> ants3$sp.div <- diversity(ants3[,6:20], index = "simpson")</pre>
```



3. Effective number of species (ENS)

Entropy and diversity

Lou Jost, Baños, Tungurahua, Ecuador (loujost@yahoo.com).

Shannon and Simpson diversity indices are misleading because they are not measured in units of species

Shannon: bits

Simpson: *probability*

Each index should be converted into the effective number of species

In a perfectly even community, ENS is equal to species richness, and always smaller than species richness in uneven communities

Effective number of species

- for species richness = S
- for Shannon index = e^H
 (exponential of Shannon entropy index)
- for Simpson index = 1/D (reciprocal of Simpson concentration index)

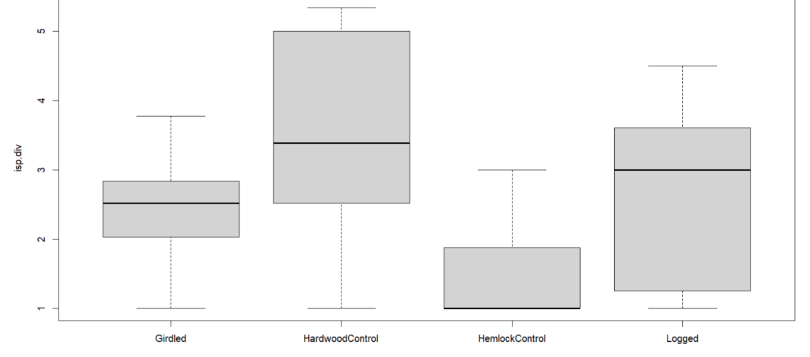
3. Effective numbers of species (ENS)

Effective number of species

 for Simpson index = 1/D (reciprocal of Simpson concentration index)

```
R package
vegan
```

```
> # Effective numbers of species - Inverse Simpson index
> diversity(ants3[,6:20], index = "invsimpson")
[1] 3.600000 3.000000 1.0000000 1.373984 3.769231 3.000000 1.000000 2.373626 1.000000 2.666667
[11] 1.000000 1.090722 3.769231 5.000000 3.000000 2.053521 2.666667 2.666667 2.000000 2.064220
[21] 4.500000 3.600000 1.142132 1.000000 3.000000 1.000000 5.000000 5.333333 1.000000 2.976517
> ants3$isp.div <- diversity(ants3[,6:20], index = "invsimpson")</pre>
```



4. Hill numbers

Several common diversity metrics can be united into a single family of generalized entropy metrics

- Parameterized by variable q

$$^qD = \left(\sum_{i=1}^S p_i^{\ q} \right)^{\frac{1}{(1-q)}}$$
 Iill numbers

For q = 0, 1 and 2 (also noted as N_0 , N_1 and N_2):

$${}^{0}D = S_{\text{(species richness)}}$$

$$^{1}D = e^{H}_{\text{(exponential of Shannon entropy)}}$$

$$^2D = \frac{1}{D}_{\text{ (reciprocal of Simpson index)}}$$

q = 0; species richness – counts all species equally without considering relative abundance

q = 1; Shannon – ENS, each species is weighted in proportion to its abundance

q = 2; Inverse Simpson – ENS, disproportionately favors abundant species

DIVERSITY AND EVENNESS: A UNIFYING NOTATION AND ITS CONSEQUENCES¹

 $M. O. Hull^2$

School of Plant Biology, University College of North Wales, Bangor, Caernarvonshire, Wales

4. Hill numbers

```
> hill_taxa(ants3[,6:20], q = 0, MARGIN = 1)
[1] 4 3 1 3 4 3 1 3 1 3 1 2 4 5 3 5 3 3 2 4 5 4 2 1 3 1 5 6 1 6
> hill_taxa(ants3[,6:20], q = 1, MARGIN = 1)
[1] 3.779763 3.000000 1.0000000 1.709090 3.864313 3.000000 1.0000000 2.583758 1.0000000 2.828427
[11] 1.000000 1.195835 3.864313 5.000000 3.000000 2.771903 2.828427 2.828427 2.0000000 2.686272
[21] 4.762203 3.779763 1.277532 1.0000000 3.0000000 1.0000000 5.0000000 5.656854 1.0000000 3.739531
> hill_taxa(ants3[,6:20], q = 2, MARGIN = 1)
[1] 3.600000 3.0000000 1.0000000 1.373984 3.769231 3.0000000 1.0000000 2.373626 1.0000000 2.666667
[11] 1.0000000 1.090722 3.769231 5.0000000 3.0000000 2.053521 2.6666667 2.6666667 2.0000000 2.976517
```

```
q = 0; species richness
```

q = 1; Shannon

q = 2; Inverse Simpson

R package

hillR

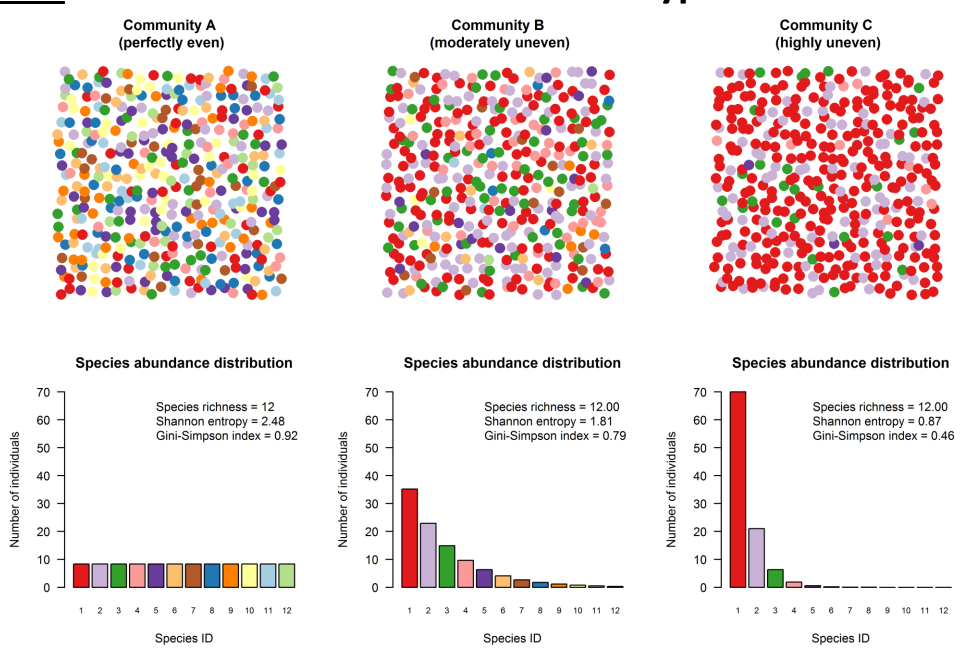
Methods in Ecology and Evolution 2016, 7, 1451–1456 doi: 10.1111/2041-210X.12613

APPLICATION

iNEXT: an R package for rarefaction and extrapolation of species diversity (Hill numbers)

T. C. Hsieh, K. H. Ma and Anne Chao*

Evenness: the relative abundance of each type



David Zelený

Species evenness

Measure of the relative abundance of species

Commonness vs rarity of species

Pielou's J – calculated as a ratio of the Shannon index to the maximum Shannon index (where all species have the same relative abundance) - Ranges from 0-1

Equitability – calculated as Simpson ENS (or inverse Simpson) divided by the observed number of species

Shannon's evenness

$$J = \frac{H}{H_{\text{max}}} = \frac{H}{\log S}$$

Simpson's evenness

$$equitability = \frac{\frac{1}{D}}{S} = \frac{1}{DS}$$

Species evenness

0.7361257

Measure of the relative abundance of species

30 0.4960861

```
> calcDiv(ants3[,6:20], type = "PielouEven")
                                              > calcDiv(ants3[,6:20], type = "HillEven", q = 2)
   PielouEven
                                                  HillEven
    0.9591479
                                                 0.9000000
    1.0000000
                                                 1.0000000
                                                                        calcDiv(ants3[,6:20], type = "PielouEven")
          NaN
                                                 1.0000000
                                                                        calcDiv(ants3[,6:20], type = "HillEven", q = 1)
    0.4878528
                                                 0.4579946
                                                                        calcDiv(ants3[,6:20], type = "HillEven", q = 2)
    0.9751060
                                                 0.9423077
    1.0000000
                                                 1.0000000
                                                 1.0000000
          NaN
    0.8640401
                                                 0.7912088
          NaN
                                              9 1.0000000
    0.9463946
                                              10 0.8888889
11
          NaN
                                              11 1.0000000
    0.2580187
                                              12 0.5453608
   0.9751060
                                              13 0.9423077
14 1.0000000
                                              14 1.0000000
   1.0000000
                                              15 1.0000000
   0.6334721
                                              16 0.4107042
   0.9463946
                                              17 0.8888889
   0.9463946
                                              18 0.8888889
  1.0000000
                                              19 1.0000000
   0.7128026
                                               20 0.5160550
   0.9697239
                                               21 0.9000000
22 0.9591479
                                               22 0.9000000
   0.3533593
                                               23 0.5710660
24
          NaN
                                               24 1.0000000
   1.0000000
                                               25 1.0000000
26
          NaN
                                               26 1.0000000
   1.0000000
                                               27 1.0000000
                                                                             Girdled
                                                                                             HardwoodControl
                                                                                                                HemlockControl
                                                                                                                                     Logged
    0.9671320
                                               28 0.8888889
                                                                                                        treatment
                                               29 1.0000000
          NaN
```

R package chemodiv

Although understanding taxonomic metrics are important, there are limitations to these approaches

Context dependencies limit comparison among studies

Spatial comparisons

Other dimensions of biological diversity that are not reflected in taxonomic approaches

- 1) Genetic diversity
- 2) Phylogenetic diversity
- 3) Functional diversity