Contrasting Diversity Values: Statistical Inferences Based on Overlapping Confidence Intervals

Ian MacGregor-Fors¹*, Mark E. Payton²

1 Red de Ambiente y Sustentabilidad, Instituto de Ecología, A.C., Xalapa, Veracruz, México, 2 Department of Statistics, Oklahoma State University, Stillwater, Oklahoma, United States of America

Abstract

Ecologists often contrast diversity (species richness and abundances) using tests for comparing means or indices. However, many popular software applications do not support performing standard inferential statistics for estimates of species richness and/or density. In this study we simulated the behavior of asymmetric log-normal confidence intervals and determined an interval level that mimics statistical tests with $P(\alpha) = 0.05$ when confidence intervals from two distributions do not overlap. Our results show that 84% confidence intervals robustly mimic 0.05 statistical tests for asymmetric confidence intervals, as has been demonstrated for symmetric ones in the past. Finally, we provide detailed user-guides for calculating 84% confidence intervals in two of the most robust and highly-used freeware related to diversity measurements for wildlife (i.e., EstimateS, Distance).

Citation: MacGregor-Fors I, Payton ME (2013) Contrasting Diversity Values: Statistical Inferences Based on Overlapping Confidence Intervals. PLoS ONE 8(2): e56794. doi:10.1371/journal.pone.0056794

Editor: Christopher Quince, University of Glasgow, United Kingdom

Received July 9, 2012; Accepted January 14, 2013; Published February 20, 2013

Copyright: © 2013 MacGregor-Fors, Payton. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: The authors have no support or funding to report.

Competing Interests: The authors have declared that no competing interests exist.

* E-mail: ian.macgregor@inecol.edu.mx

Introduction

Measuring biodiversity is one of the major goals of ecologists around the world [1]. As suggested by Hubbell [2], biodiversity can be summarized by the species richness and relative abundances of a community in a given space and time. For decades, ecologists have used many different methods to calculate and contrast species richness, relative abundances, and/or diversity values. Most simply, ecologists often contrast species richness and abundances relative to sampling effort among different conditions using tests for comparing means (e.g., ANOVA, Kruskal-Wallis; [3–5]). Also, many indices have been developed to measure species richness and diversity (see Moreno [1,6–8] for further details). However, many popular software applications do not support performing standard inferential statistics for estimates of diversity (e.g., species richness, density).

Recently, the use of two methods for quantifying species richness and individual densities have become very popular due to their robustness: (1) rarefaction curves produced by randomly resampling the pool of total individuals or sampling units, plotting the estimated number of species in relation to a given number of individuals or sampling units [9–11], and (2) distance-sampling calculation of densities (number of individuals per area unit - e.g., hectares, square kilometers), calculated based on the probability of detection of individuals at increasing distances from the observer and the size of the successfully surveyed area [8]. Both methods can be calculated using freeware. Rarefaction curves can be generated using the output from the software EstimateS [12], which computes the expected number of species as a function number of accumulated samples (sample-based rarefaction, denoted Sobs [Mao Tao] in EstimateS) with symmetric 95% confidence intervals (Sobs 95% CI Upper and Lower Bounds). Densities can be calculated using the software Distance [13], for which asymmetric 95% confidence intervals, based on assuming the distributions of the density estimate is log-normal, are output as a default by the program.

As software programs such as EstimateS and Distance output results that cannot be contrasted directly though inferential statistics, degree of overlap between confidence intervals has been proposed to assess statistical differences [14]. Such comparisons allow testing null hypotheses regarding different environmental conditions (e.g., habitats, treatments). Although other approaches to hypothesis testing for Distance have been shown to contrast density values effectively (e.g., ANOVA, $t$-tests), they often require experience using sophisticated processes in statistical packages.

As demonstrated by Payton et al. [14], when comparing overlapping 95% confidence intervals of independent treatments with similar standard errors, non-overlapping confidence intervals represent significant differences in expectations with extremely low probabilities of Type I error ($\alpha < 0.01$), while no statistical inferences can be drawn with certainty if confidence intervals overlap but are not coincident. However, Payton et al. [14] showed that comparing 83–84% confidence intervals, instead of 95%, represents statistical tests with an $\alpha$ level of 0.05 (Fig. 1), the conventional criterion of significance for biological and ecological analyses [15].

As the 83–84% rule has previously been demonstrated only for normally distributed confidence intervals, in this study we simulated how asymmetric log-normal confidence intervals behave and determined a confidence interval level for mimicking two-sample statistical tests with $\alpha = 0.05$. As the log-normal distribution is a normal distribution on the log-scale, we predicted that the 83–84% rule should also apply to asymmetric log-normal confidence intervals. We also describe how to calculate different percentage
Results

For almost all of the scenarios contrasting samples with different means, the 84% confidence intervals provided overlap probability that best mimicked a two-tailed two population test with a 0.05 error rate. To mimic a 0.01 test, 94% confidence intervals would appear to be the proper choice. Confidence intervals at the 76% level best mimic a test with a 0.10 error rate (Tables 1, 2).

Discussion

As predicted, our results show that comparing the overlap, or lack of it, between 84% asymmetric confidence intervals pertaining to different means mimics 0.05 tests surprisingly well (Fig. 1, 2). Thus, this study provides empirical evidence that the 84% rule is suitable for mimicking 0.05 statistical tests for both symmetric and asymmetric confidence intervals. However, we did not explore the statistical power of the method (regarding Type II errors), since the primary concern of this paper was to create a process that best mimicked an alpha-level test, and the use of overlapping 84% confidence intervals for this method would be more powerful, by definition, than using 95% intervals. Assessing power for this situation would involve constructing distributions with different means (and, by virtue of the nature of the log-normal distribution, different variances) and assessing the ability of the method to detect differences in overlapping confidence intervals with different means. Though our results have been demonstrated effective only for normal symmetric intervals and for log-normal asymmetric intervals, we believe that the 84% rule for mimicking 0.05 tests with overlapping confidence intervals might work effectively for other distributions. For example, comparing 84% confidence intervals for species estimation comparisons using widely used non-parametric estimators (e.g., Chao1, Chao2, ICE, ACE, Jackknife, Bootstrap), could mimic 0.05 tests. However, it remains to be tested.

Materials and Methods

Simulations for Mimicking Pairwise Tests Based on Asymmetric Confidence Intervals

We performed simulations to establish the confidence intervals at which P<0.01, 0.05, and 0.10 Type I error was achieved, mimicking pairwise tests with PC SAS [16]. In order to explore how the proposed method behaves for various types of log-normal distributions, we created several combinations of the two parameters of the log-normal distribution (μ and σ). Specifically, we created 48 different log-normal distributions by utilizing 6 different levels for μ and 8 different levels for σ in an effort to cover a variety of different distributions. For the purposes of these simulations, we generated samples from parent populations which were generated by assuming different means and corresponding standard errors, which are functions of the parameters utilized to create these parent populations. Thus, as we assessed the behavior of asymmetric confidence intervals, we calculated a confidence interval for each of two samples drawn from the same population, each with alpha values varying from 0.05 to 0.25, at 0.01 increments. We calculated 10,000 iterations of each simulation scenario, including populations with different means extracted from the same parent populations. For each iteration, we calculated 0.75% to 0.95% confidence intervals in 1% increments, and we used this series of confidence intervals to determine the proportion of times the simulated confidence intervals overlap for each nominal level of confidence. Note that the log-normal distribution’s coefficient of variation is a function of σ only [17], so changing the mean of the distribution changes, by definition, the variance also.
In order to generate 84% confidence intervals for rarefaction analyses, the standard deviation of the observed species (Mao Tao SD) from the output file from EstimateS is needed. As standard deviations equal standard errors in EstimateS because infinite degrees of freedom are assumed in the calculation of Mao Tao SD, the latter must be multiplied by 1.372, the quantile (normal curve z-score) corresponding to two-sided intervals of 84% probabilities, with alpha = 0.16, and cumulative probabilities of 0.08 and 0.92. For example, if Mao Tao SD = 5.55, for example, 84% confidence intervals for that specific value of Mao Tao SD, which can vary in relation to the number of accumulated computed individuals in a rarefaction plot, are equal to the average value 6.76.

As the Distance program can calculate user-selected levels for confidence intervals (default = 0.95) for distance-sampling density calculations, setting the confidence interval limits solves the issue. To accomplish this, go the “Analyses” button on the toolbar, select “Analysis details” and a new window will appear. Finally, select the “Misc” tab and modify the default value for confidence intervals (i.e., 95) to 84. Results output from the Distance program will now include 84% confidence intervals.

Wildlife species richness and density measurements of ecosystems are imperative in order to concentrate conservation actions in highly biodiverse areas [1]. In this paper, we demonstrated that the 84% rule mimics 0.05 pairwise statistical tests for both symmetric and asymmetric confidence intervals, with detailed users’ guides for calculating 84% confidence intervals in two of the most robust and highly-used freeware applications related to biodiversity (i.e., EstimateS, Distance). Thus, we encourage ecologists to use these programs to calculate species richness and individual

| μ  | σ  | 2  | Mean  | Variance | Size of CIs (%) | Average overlap |
|----|----|----|-------|----------|----------------|-----------------|
| 4.5| 0.0005| 90.04 | 4.05 | 84 | 0.9541 |

Results from simulations including 10,000 iterations.

doi:10.1371/journal.pone.0056794.t002

Figure 2. Comparison of the use of 95% and 84% confidence intervals in three replicates of our simulations. For this representative example, the data were created from a log-normal population with a mean of 90.2 and variance of 32.6. In case 1, the both sets of intervals overlap, both suggesting that no significant (NS) differences exist. Note, however, that the 95% confidence intervals will yield an error rate of less than 1%, while the 84% confidence intervals better mimic a 0.05 level test. In case 2, 95% confidence intervals slightly overlap, while 84% ones do not. For this situation, these two approaches would lead to different conclusions: (a) significant differences (*) when considering 84% confidence intervals, and (b) no statistical differences can be inferred using 95% confidence intervals (?). In case 3, none of the sets of intervals overlap, both suggesting that significant differences exist. Note, however, that statistical differences using 95% confidence intervals are assumed with an error rate of less than 1%, while that of 84% confidence intervals better mimic a 0.05 level test.

doi:10.1371/journal.pone.0056794.g002

In order to generate 84% confidence intervals for rarefaction analyses, the standard deviation of the observed species (Mao Tao SD) from the output file from EstimateS is needed. As standard deviations equal standard errors in EstimateS because infinite degrees of freedom are assumed in the calculation of Mao Tao SD, the latter must be multiplied by 1.372, the quantile (normal curve z-score) corresponding to two-sided intervals of 84% probabilities, with alpha = 0.16, and cumulative probabilities of 0.08 and 0.92. For example, if Mao Tao SD = 5.55, for example, 84% confidence intervals for that specific value of Mao Tao SD, which can vary in relation to the number of accumulated computed individuals in a rarefaction plot, are equal to the average value ±7.61.

As the Distance program can calculate user-selected levels for confidence intervals (default = 0.95) for distance-sampling density calculations, setting the confidence interval limits solves the issue. To accomplish this, go the “Analyses” button on the toolbar, select “Analysis details” and a new window will appear. Finally, select the “Misc” tab and modify the default value for confidence intervals (i.e., 95) to 84. Results output from the Distance program will now include 84% confidence intervals.

Wildlife species richness and density measurements of ecosystems are imperative in order to concentrate conservation actions in highly biodiverse areas [1]. In this paper, we demonstrated that the 84% rule mimics 0.05 pairwise statistical tests for both symmetric and asymmetric confidence intervals, with detailed users’ guides for calculating 84% confidence intervals in two of the most robust and highly-used freeware applications related to biodiversity (i.e., EstimateS, Distance). Thus, we encourage ecologists to use these programs to calculate species richness and individual
density statistical expectations, applying this easy-to-use overlapping confidence interval method when making statistical inferences, which represents an alternative to the use of diversity indices.

Acknowledgments

We are very thankful to Len Thomas, Robert K. Colwell, Anne Chao, Javier Quesada, and Federico Escobar for their valuable comments and ideas.

References

1. Magurran AE (2004) Measuring Biological Diversity. Oxford: Blackwell Publishing.
2. Hubbell SP (2001) The Unified Neutral Theory of Biodiversity and Biogeography. Princeton: Princeton University Press.
3. Fernández-Juricic E (2001) Avian spatial segregation at edges and interiors of urban parks in Madrid, Spain. Biodivers Conserv 10: 13031–316.
4. Gaines WL, Haggard M, Lehmkahl JF, Lyons A L, Harrod RJ (2007) Short-term response of land birds to Ponderosa Pine restoration. Rest Ecol 15: 670–678.
5. Moreno C, Rodríguez P (2010) A consistent terminology for quantifying species diversity? Oecologia 163: 279–282.
6. Moreno CE (2001) Métodos para Medir la Biodiversidad. Zaragoza: M&T-Manuales y Tesis SEA.
7. Magurran AE, McGill BJ (2011) Biological Diversity: Frontiers in Measuring Biodiversity. New York: Oxford University Press.
8. Buckland ST, Stouden EC, Magurran AE, Illian J, Newson JE (2011) The geometric mean of relative abundance indices: A biodiversity measure with a difference. Ecosphere 2: 100.
9. Gotelli NJ, Colwell RK (2001) Quantifying biodiversity: Procedures and pitfalls in the measurement and comparison of species richness. Ecol Lett 4: 379–391.

Author Contributions

Conceived and designed the experiments: IMF MEP. Performed the experiments: MEP. Analyzed the data: MEP IMF. Contributed reagents/materials/analysis tools: MEP IMF. Wrote the paper: IMF MEP.

10. Gotelli NJ, Colwell RK (2011) Estimating species richness. In: Magurran AE, McGill BJ, editors. Frontiers in Measuring Biodiversity. New York: Oxford University Press. pp 39–54.
11. Colwell RK, Chao A, Gotelli NJ, Lin SV, Mao CX, et al. (2012) Models and estimators linking individual-based and sample-based rarefaction, extrapolation, and comparison of assemblages. J Plant Ecol 5: 3–21.
12. Colwell RK (2011) EstimateS: Statistical estimation of species richness and shared species from samples, Version 9. Available: http://viceroy.eeb.uconn.edu/estimates. Accessed 2012 Sep 7.
13. Thomas L, Buckland ST, Rexstad EA, Laake LJ, Strindberg S, et al. (2010) Distance software: Design and analysis of distance sampling surveys for estimating population size. J Appl Ecol 47: 5–14.
14. Payton ME, Greenstone MH, Schenker N (2003) Overlapping confidence intervals or standard error intervals: What do they mean in terms of statistical significance? J Insect Sci 3: 34.
15. Sokal RR, Rohlf FJ (1995) Biometry: The Principles and Practice of Statistics in Biology Research. New York: Freeman.
16. SAS (2008) PC SAS Version 9.2. Cary, NC: SAS Institute.
17. Limpert E, Stahel WA, Abbe M (2001) Log-normal distributions across the science: Keys and clues. BioScience 51: 341–352.