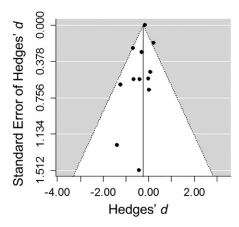




## Reply to Salkeld et al.: Diversity-disease patterns are robust to study design, selection criteria, and publication bias

Recently, we provided broad evidence for a negative association between community diversity and the abundance of human and wildlife parasites (1), supporting the dilution effect hypothesis (2). Salkeld et al. (3) point out that our meta-analysis reached a different conclusion than their previous meta-analysis on the dilution effect (4), which focused solely on 13 effect sizes from six species of human parasites. Salkeld et al. (3) argue that their smaller sample size is justified because only field studies capture the ecological realism necessary to test the dilution effect hypothesis. However, field studies can have confounding variables that can generate false-positive or false-negative relationships between diversity and disease. Although laboratory studies often more confidently capture cause-effect relationships, they can be contrived. Hence, given these costs and benefits of field and laboratory studies, we included both in our meta-analysis and found that each strongly supported the dilution



**Fig. 1.** Funnel plot for supplementary analysis of mean effect sizes for the relationship between biodiversity and parasite abundance for 12 human zoonotic parasites. There is no evidence for funnel plot asymmetry, a potential indicator of publication bias, using either the rank correlation test or Egger's regression test implemented in the metafor package in R (6).

effect hypothesis. Nevertheless, we reanalyzed the data in our publicly available dataset using the inclusion criteria (i.e., field studies on human parasites at certain scales) of Salkeld et al. (4) and, once again, found a significant negative relationship between biodiversity and the abundance of human parasites (random effects meta-analysis: n = 22 effect sizes and 12 species as a random effect, mean Hedges' d = -0.75, P = 0.018). Hence, the difference between the marginally nonsignificant finding in the meta-analysis of Salkeld et al. (mean Fisher's Z = -0.23, 95% confidence interval: -0.470 to 0.008) (4) and our significant finding can be explained by the increased statistical power associated with the inclusion of nine more recent studies on six human parasites. These results further support the robustness of the negative relationship between biodiversity and infectious disease using our dataset.

Salkeld et al. (3) also suggest that publication bias might influence the biodiversity-disease literature, and we agree that this bias is possible. We emphasized (1) that we did not test for publication bias because the currently available diagnostic tools are invalidated when effect sizes share some underlying replicates, come from the same larger experiment, or are on the same species [i.e., are not independent (5)]. These sources of dependence are also present in the dataset of Salkeld et al. (4), compromising their original tests of publication bias. Despite this caveat, we here explore for signs of bias using the weighted means and variances of Hedges' d for each human parasite species, again using the selection criteria of Salkeld et al. (n = 12) (4). Neither the funnel plot (Fig. 1) nor diagnostic statistical tests suggest publication bias (rank correlation test: Kendall's  $\tau = -0.24$ , P = 0.31; Egger's regression test: z = -1.05, P = 0.29) (6).

We reiterate that the evidence supporting the dilution effect hypothesis is robust to study design, meta-analysis selection criteria, and publication bias. However, the dilution effect is a heterogeneous phenomenon; therefore, in some instances, increasing biodiversity could increase rather than decrease disease (1). Hence, when the goal is to manage a known problematic disease, obtaining system-specific information might improve the success of disease control over blindly conserving biodiversity. In contrast, if the goal is to minimize disease emergence/abundance without species-specific knowledge, conserving biodiversity may be a useful strategy.

David J. Civitello<sup>a, I</sup>, Jeremy Cohen<sup>a</sup>, Hiba Fatima<sup>b</sup>, Neal T. Halstead<sup>a</sup>, Taegan A. McMahon<sup>c</sup>, C. Nicole Ortega<sup>a</sup>, Erin L. Sauer<sup>a</sup>, Suzanne Young<sup>a</sup>, and Jason R. Rohr<sup>a</sup>

<sup>a</sup>Department of Integrative Biology, University of South Florida, Tampa, FL 33620; <sup>b</sup>Global Health Institute, Duke University, Raleigh, NC 22708; and <sup>c</sup>Department of Biology, University of Tampa, Tampa, FL 33606

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The authors declare no conflict of interest.

<sup>1</sup>To whom correspondence should be addressed. Email: civitello@