Effect Sizes for Single-Case Research

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Effect sizes: numerical indices quantifying magnitude and direction of an intervention effect (functional relationship) on a scale that is comparable across cases/studies.

Why does single-case research need effect size measures?

1. Synthesizing the evidence base for an intervention/practice using meta-analysis
2. Examining heterogeneity of effects (for whom and under what conditions is a treatment effective?)
3. Studying the process of evidence production (meta-science)
WHAT KIND OF EFFECT SIZES DO WE NEED?

- Comparability *across a set of cases/studies to be synthesized*

- Studies might use different measurement procedures to assess a common outcome construct (dependent variable)
  - E.g., continuous recording, momentary time sampling, partial interval recording of a behavior

- Studies might use different research designs to evaluate a common intervention (functional relationship)
  - E.g., multiple baseline, treatment reversal, between-case experimental designs
BETWEEN-CASE STANDARDIZED MEAN DIFFERENCES (BC-SMD)

- **SMD** is the most familiar effect size index in between-case experimental research
  - Difference in mean outcomes (treated – not treated)
  - Scaled by cross-sectional SD of outcome

- **BC-SMDs** (Hedges, Pustejovsky, & Shadish, 2012, 2013; Pustejovsky, Hedges, & Shadish, 2014; Shadish, Hedges, & Pustejovsky, 2014) estimate *the same effect size parameter as in a between-case experimental design*, using data from a single-case design.

- Translate single-case results into terms that are familiar for between-case researchers (Shadish, Hedges, Horner, & Odom, 2015)
Methods currently available for:

- Across-participant multiple baseline/multiple probe design
- Treatment reversal design with replication across cases
- Requires studies with at least 3 participants, outcomes measured on a common scale
TOOLS FOR CALCULATING BC-SMDS

- SPSS macro (Shadish & Marso, 2015)
- R package `scdhlm` available on CRAN (Pustejovsky, 2016)
- Shiny web-app now available (https://jepusto.shinyapps.io/scdhlm/)
A demonstration of how to do a meta-analysis that combines single-case designs with between-groups experiments: The effects of choice making on challenging behaviors performed by people with disabilities

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**ABSTRACT**

*Objective:* This paper demonstrates how to conduct a meta-analysis that includes both between-group and single-case design (SCD) studies. The example studies whether choice-making interventions decrease challenging behaviors performed by people with disabilities. *Methods:* We used a between-case $d$-statistic to conduct a meta-analysis of 15 between-group and SCD studies of 70 people with a disability, who received a choice intervention or control. We used robust variance estimation to adjust for dependencies caused by multiple effect sizes per study, and conducted moderator, sensitivity, influence, and publication bias analyses. *Results:* The random-effects average was $d = 1.02$ (standard error of 0.168), so the 95% confidence interval (CI) suggests choice-making reduces challenging behaviors by 0.65 to 1.38 standard deviations. Studies that provided choice training produced a significantly larger intervention effect. *Conclusion:* Choice-making reduces challenging behaviors performed by people with disabilities.

**KEYWORDS**

Meta-analysis; challenging behaviors; robust variance estimation; choice; disabilities
Single-case experimental design yielded an effect estimate corresponding to a randomized controlled trial

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Abstract

Objectives: We reanalyzed data from a previous randomized crossover design that administered high or low doses of intravenous immunoglobulin (IgG) to 12 patients with hypogammaglobulinaemia over 12 time points, with crossover after time 6. The objective was to see if results corresponded when analyzed as a set of single-case experimental designs vs. as a usual randomized controlled trial (RCT).

Study Design and Settings: Two blinded statisticians independently analyzed results. One analyzed the RCT comparing mean outcomes of group A (high dose IgG) to group B (low dose IgG) at the usual trial end point (time 6 in this case). The other analyzed all 12 time points for the group B patients as six single-case experimental designs analyzed together in a Bayesian nonlinear framework.

Results: In the randomized trial, group A ($M = 794.93; \text{standard deviation} (SD) = 90.48$) had significantly higher serum IgG levels at time six than group B ($M = 283.89; SD = 71.10$) ($t = 10.88; df = 10; P < 0.001$), yielding a mean difference of $M_D = 511.05$ [standard error (SE) = 46.98]. For the single-case experimental designs, the effect from an intrinsically nonlinear regression was also significant and comparable in size with overlapping confidence intervals: $M_D = 495.00, SE = 54.41, \text{and } r = 495.00/54.41 = 9.10$. Subsequent exploratory analyses indicated that how trend was modeled made a difference to these conclusions.

Conclusions: The results of single-case experimental designs accurately approximated results from an RCT, although more work is needed to understand the conditions under which this holds. © 2016 Elsevier Inc. All rights reserved.
WITHIN-CASE EFFECT SIZES

- Within-case = characterize magnitude of functional relationship separately for each case in a study.

- Many different indices have been proposed
  - parametric measures (e.g., within-case standardized mean difference)
  - non-overlap measures (e.g., PND, IRD, NAP, Tau-U).

- Need for effect sizes that are on comparable scale across cases/studies that use different outcome measurement procedures.
  - Outcome recording system
  - Observation session length
  - Phase lengths
# Measurement Procedures in Five Systematic Reviews of Single-Case Research

<table>
<thead>
<tr>
<th>Procedure</th>
<th>Object play (Barton)</th>
<th>Functional behavior assessment (Gage)</th>
<th>Antecedent social skills (Ledford)</th>
<th>Group contingencies (Maggin)</th>
<th>Choice-making (Shogren)</th>
<th>Total</th>
</tr>
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<td>13 (7%)</td>
<td>213 (32%)</td>
<td>61 (36%)</td>
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<td>49 (7%)</td>
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<td>4 (6%)</td>
<td>9 (1%)</td>
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</tbody>
</table>
OBSERVATION SESSION LENGTHS (MIN) IN FIVE SYSTEMATIC REVIEWS OF SINGLE-CASE RESEARCH

- Choice-making (Shogren)
- Group contingencies (Maggin)
- Antecedent social skills (Ledford)
- Functional behavior assessment (Gage)
- Object play (Barton)
Log response ratios quantify functional relationships in terms of proportionate change between phases.

- Common “informal” effect size measure (Campbell & Herzinger, 2010)
- Precedents in systematic reviews of single-case research (Campbell, 2003; Kahng, Iwata, & Lewin, 2002; Marquis et al., 2000)

Letting $\mu_A, \mu_B$ denote mean levels of outcome in phases A and B, LRR is

$$\psi = \ln\left(\frac{\mu_B}{\mu_A}\right) = \ln(\mu_B) - \ln(\mu_A)$$

Useful for behavioral outcomes measured by direct observation (Pustejovsky, 2015)
- Magnitude remains stable when outcomes are measured using different procedures.
- Under certain conditions, also comparable across dimensional constructs.
Further development of within-case log response ratios
- to accommodate linear/non-linear time trends
- Estimation in the presence of auto-correlated repeated measurements

Choice of effect size metric depends on characteristics of set of studies to be synthesized.

Publication/outcome reporting bias represents a major outstanding challenge for synthesis of single-case research.
A SURVEY OF PUBLICATION PRACTICES OF SINGLE-CASE DESIGN RESEARCHERS WHEN TREATMENTS HAVE SMALL OR LARGE EFFECTS

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The published literature often underrepresents studies that do not find evidence for a treatment effect; this is often called publication bias. Literature reviews that fail to include such studies may overestimate the size of an effect. Only a few studies have examined publication bias in single-case design (SCD) research, but those studies suggest that publication bias may occur. This study surveyed SCD researchers about publication preferences in response to simulated SCD results that show a range of small to large effects. Results suggest that SCD researchers are more likely to submit manuscripts that show large effects for publication and are more likely to recommend acceptance of manuscripts that show large effects when they act as a reviewer. A non-trivial minority of SCD researchers (4% to 15%) would drop 1 or 2 cases from the study if the effect size is small and then submit for publication. This article ends with a discussion of implications for publication practices in SCD research.

Key words: single-case design, publication bias, effect size


