Review Article

Trial Characteristics and Appropriateness of Statistical Methods Applied for Design and Analysis of Randomized School-Based Studies Addressing Weight-Related Issues: A Literature Review

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Objective. To evaluate whether clustering effects, often quantified by the intraclass correlation coefficient (ICC), were appropriately accounted for in design and analysis of school-based trials. Methods. We searched PubMed and extracted variables concerning study characteristics, power analysis, ICC use for power analysis, applied statistical models, and the report of the ICC estimated from the observed data. Results. N = 263 papers were identified, and N = 121 papers were included for evaluation. Overall, only a minority (21.5%) of studies incorporated ICC values for power analysis, fewer studies (8.3%) reported the estimated ICC, and 68.6% of studies applied appropriate multilevel models. A greater proportion of studies applied the appropriate models during the past five years (2013–2017) compared to the prior years (74.1% versus 63.5%, p = 0.176). Significantly associated with application of appropriate models were a larger number of schools (p = 0.030), a larger sample size (p = 0.002), longer follow-up (p = 0.014), and randomization at a cluster level (p < 0.001) and so were studies that incorporated the ICC into power analysis (p = 0.016) and reported the estimated ICC (p = 0.030). Conclusion. Although application of appropriate models has increased over the years, consideration of clustering effects in power analysis has been inadequate, as has report of estimated ICC. To increase rigor, future school-based trials should address these issues at both the design and analysis stages.

1. Introduction

Pediatric and adolescent obesity is a global concern as the range of its health consequence includes cardiovascular diseases, diabetes, poor quality of life, disability, mental health problems, and even adulthood mortality [1–5]. Despite this concern, the prevalence of pediatric and adolescent obesity has not decreased in the United States (US) or globally over the past several decades [6, 7]. In addition, financial and care burdens for preventing and treating pediatric obesity are substantial at both the individual or family level and societal or governmental level [8]. As such, countless trials have been conducted over the world to address pediatric and adolescent obesity and prevention, treatment, or diet guidelines based on evidence collected from the findings of those trials have been published [9, 10]. Many opine that schools represent “key settings” or “ideal settings” for obesity prevention or intervention, and
numorous school-based studies have been conducted world-
wide [11, 12]. The scope of interventions in school-based trials
is broad ranging from education to health behaviors focusing
on nutrition and physical activities [13–15]. However, the
appropriate evaluation of intervention effectiveness critically
hinges on proper trial design and statistical analysis of school-
based trial data. Importantly, the data from school-based trials
naturally form multiple levels of hierarchy. For example, when
students are to be followed up multiple times for outcome
measurements during the course of a study, repeatedly
measured outcomes are nested within students who are in turn
nested within schools. This hierarchical nature forms a three-
level data structure so that clustering effects, also known as
design effects, of outcome data at the school and student levels
should be taken into account not only at the design stage but
also at the analysis stage.

The critical issue is that required sample sizes are likely to be
underestimated if such clustering effects are not taken into
account [16–20]. This is especially so when the interventions
are assigned at the highest level of data hierarchy. Likewise, the
standard errors of estimated intervention effects at the analytic
stage are smaller than what it should be and so are $p$ values,
likely increasing the type I error rate [21]. Anecdotally, errors in
neglecting such clustering seem not uncommon [22].

The primary aim of this review is to evaluate statistical
methods applied to published school-based randomized
trials addressing weight issues. The evaluation is focused on
assessing appropriateness of power analysis and applied
statistical methods as to whether clustering effects often
represented by correlations of subject outcomes within
schools, which is also known as the intracluster (or intra-
class) correlation coefficient (ICC) [23], are properly
accounted for in the design and analysis of school-based
trials. Specifically, we aim to examine (1) whether study
characteristics are different between the past five years
(2013–2017) and the prior years (1995–2012); number of schools; analytic
sample sizes of individuals, which are the size of the comple-
ters at the final follow-up when available; randomization
level (cluster versus individual); length of follow-up in months
(<12 versus ≥12 months); number of repeated measure-
ments including baseline (2 versus ≥2); location of the trial (the
US versus others); and types of schools (preschool, elementary/
primary, middle, and high school, and college).

As per the randomization level, the “cluster” classification
category includes any higher level units in which all subjects
received the identical interventions or treatments. Therefore,
the cluster unit could be schools, classes, health centers,
communities, camps, and so on. Nevertheless, we extracted
the number of involved schools, instead of the number of the
various randomized cluster units, for the number of schools
variable. If randomization occurred at the individual/student
level within schools or other cluster units, we classified the
randomization level for such trials as “individual.” When
multiple types of schools are involved in a trial, we classified
such cases into the school type of the lowest level. For in-
stance, if a trial involved preschool and primary schools, then
this case was classified as “preschool” for the school type.

With respect to the presence of power analysis and
appropriateness of applied statistical methods, we extracted
the report of power analysis (yes versus no); consideration of
the intracluster correlation coefficient (ICC) for power
analysis (yes versus no); primary statistical analysis models
(multilevel versus no); and the report of the ICC estimated
from statistical analysis (yes versus no). If the description
of power analysis or sample size determination referred to
a protocol paper and was not provided in the text to an

2. Methods

2.1. Search Strategy. We searched PubMed for papers
meeting the following inclusion criteria:

(i) Published in journals indexed in PubMed
(ii) From the PubMed inception year
(iii) Randomized school-based trials
(iv) Written in English
(v) Outcomes involving weight-related issues
(vi) Subjects of age younger than 25 years that include
young adults.

To this end, we applied a Boolean algebraic combination of
MeSH (Medical Subject Headings) as follows: (“Body
Weight”[MeSH] OR “Body Mass Index”[MeSH]) AND
“Schools”[MeSH] AND (Randomized Controlled Trial[ptyp]
AND English[lang] AND (“adolescent”[MeSH Terms] OR
“child”[MeSH Terms:noexp] OR “child, preschool”[MeSH
Terms] OR “young adult”[MeSH Terms])). With this search
strategy, we identified a total of 263 papers as of December 8,
2017, and the earliest publication year was 1995.

2.2. Exclusion Criteria. We retrieved full texts of all of those
papers, and two authors (MH and SN) applied the following
exclusion criteria for the review:

(i) Protocol paper.
(ii) Not a randomized study.
(iii) Not a school-based intervention.
(iv) Analysis of subjects of age beyond the range.
(v) Outcome is not related to weight issues.
(vi) Baseline or subgroup analysis of a parent study.
(vii) Single-school trials.

When the two raters’ ratings were not concordant due to
unclear descriptions in the text regarding study design
parameters or analytic methods, the rating was resolved by
consensus between them.

2.3. Extracted Variables and Classifications. We extracted the
following study characteristic variables from the included
studies: the publication year (past 5 years (2013–2017) versus
the prior years (1995–2012)); number of schools; analytic
sample sizes of individuals, which are the size of the com-
toters at the final follow-up when available; randomization
level (cluster versus individual); length of follow-up in months
(<12 versus ≥12 months); number of repeated measure-
ments including baseline (2 versus ≥2); location of the trial (the
US versus others); and types of schools (preschool, elementary/
primary, middle, and high school, and college).

As per the randomization level, the “cluster” classification
category includes any higher level units in which all subjects
received the identical interventions or treatments. Therefore,
the cluster unit could be schools, classes, health centers,
communities, camps, and so on. Nevertheless, we extracted
the number of involved schools, instead of the number of the
various randomized cluster units, for the number of schools
variable. If randomization occurred at the individual/student
level within schools or other cluster units, we classified the
randomization level for such trials as “individual.” When
multiple types of schools are involved in a trial, we classified
such cases into the school type of the lowest level. For in-
stance, if a trial involved preschool and primary schools, then
this case was classified as “preschool” for the school type.

With respect to the presence of power analysis and
appropriateness of applied statistical methods, we extracted
the report of power analysis (yes versus no); consideration of
the intracluster correlation coefficient (ICC) for power
analysis (yes versus no); primary statistical analysis models
(multilevel versus no); and the report of the ICC estimated
from statistical analysis (yes versus no). If the description
of power analysis or sample size determination referred to
a protocol paper and was not provided in the text to an
evaluable extent, we classified such cases as no report of power analysis. As the terminologies of statistical models are not standardized across the papers, we classified the following models as “multilevel” models in which a “clustering” effect at some level of the data hierarchy had presumably been taken into account in the analysis: generalized mixed-effects model, mixed-effects model, linear mixed-effects model, multilevel mixed-effects model, hierarchical models, random-intercept models, mixed models, logistic mixed-effects models, generalized estimating equation models, and the likes. We consider the application of those multilevel analyses appropriate. Other models that do not take the clustering effects into account were classified as “no” multilevel model: for example, paired t-test, chi-square test, analysis of variance (ANOVA), analysis of covariance (ANCOVA), linear or logistic regression models, and repeated-measures AN(C)OVA at the individual level.

2.4. Statistical Analysis. Descriptive statistics are provided in terms of frequency, percentages, median, and the first (Q1) and the third (Q3) quartiles. There were three missing values for the number of schools, and studies with those missing values were excluded from analyses involving the number of schools. The follow-up length and number of measurements are analyzed as both continuous and dichotomized variables. Comparisons of study characteristics, the presence of power analysis, and appropriateness of statistical methods between the publication years 2013–2017 versus 1995–2012 are made using Wilcoxon rank-sum tests or Fisher exact tests. The same sets of these tests are also applied to testing associations of study characteristics and the presence of power analysis with application of appropriate multilevel analysis. Statistical significance is declared at \( p < 0.05 \) (two-tailed). SAS v9.4 was used for all analyses.

3. Results

3.1. Reasons for Exclusion. The application of the exclusion criteria resulted in 121 papers (Supplementary Materials (available here)) with 1995 as the earliest publication year after exclusion of 142 papers. The reasons for exclusions are displayed in Table 1. Although it is possible that a study was excluded for multiple reasons, the most common reason was due to single-school trials (34.5%) followed by protocol papers (25.4%).

| Reasons                              | N  | %    |
|--------------------------------------|----|------|
| Single-school trial                  | 49 | 34.5 |
| Protocol paper                       | 36 | 25.4 |
| Baseline or subgroup analysis of a parent study | 27 | 19.0 |
| Not a randomized trial               | 18 | 12.7 |
| Students of age beyond the age range | 7  | 4.9  |
| Not a school-based trial             | 5  | 3.5  |
| Total                                | 142|      |

Note. Although it is possible that studies could be excluded for multiple reasons, we classified the reasons in a mutually exclusive manner.

3.2. Study Characteristics between the Past Five Years and the Prior Years. Detailed results are presented in Table 2. There were 63 (52%) and 58 (48%) studies included for analysis between the years 2013–2017 versus 1995–2012, respectively. None of the listed study characteristics, except the school type, significantly differ between those time frames. We note that the initial extended CONSORT guideline statement for cluster randomized trials was published in the year 2004 [25], but we did not further categorize years into three time frames including those before 2005 due to a relatively small number of included studies during that time period (\( N = 19 \) or 7.3%). Overall, the vast majority of studies randomized interventions at a cluster level (89.3%), and the analytic sample size is relatively large with a median of 620 students. The majority (58.9%) of studies had only two measurements: one at baseline or before intervention and the other at the end of the trial/follow-up or after intervention. Most (57.0%) of the studies were conducted outside the US, and more so for the past five years compared to the prior years (63.8% versus 50.8%, \( p = 0.198 \)). The number of countries outside the US was 30 from all continents; one study involved six European countries. The primary/elementary schools most often served as experimental settings (68.6%); however, those schools were significantly less utilized during the past 5 years compared to the prior years (56.9% versus 79.4%, \( p = 0.010 \)). Perhaps for this reason, the distributions of overall types of schools are significantly different between years (\( p = 0.038 \)).

3.3. Power Analysis, ICC, and Multilevel Models between the Past Five Years and the Prior Years. Results from the comparisons of power analysis and application of multilevel models between years are presented in Table 3. A minority (43.0%) of the studies reported power analysis adequately in the text, and the proportion was not significantly different between the past five years and the prior years (48.3% versus 38.1%, \( p = 0.258 \)). However, most studies applied multilevel analytic methods (68.6%), and the proportion was greater for the past five years (74.1% versus 63.5%, \( p = 0.176 \)). About one-fifth of studies (21.5%) took the ICC into consideration for power analysis, and the past five years did not see a significant increase in this proportion. A smaller number of studies (8.3%) reported ICCs estimated from the statistical analysis of trial data, and the proportions decreased, though not significantly, during the past five years.

3.4. Association between Multilevel Analysis and Study Characteristics. As presented in Table 4, a majority of study characteristics are associated with the application of appropriate multilevel statistical models. Specifically, a larger number of schools (\( p = 0.030 \)), a larger size of the analytic sample (\( p = 0.002 \)), longer follow-up (\( p = 0.014 \)), and randomization at a cluster level (\( p < 0.001 \)) were all significantly associated with the application of appropriate models. The location of trials was not significantly associated with the appropriateness (\( p = 1.000 \)). The studies with appropriate multilevel models were more likely to have incorporated the ICC into power analysis (\( p = 0.016 \) and reported the ICC
from the analysis of trial data ($p = 0.030$). As should be the case, no studies with non-multilevel models reported ICC estimates. However, even among studies with multilevel models, only a minority (12.1%) reported ICC estimates.

## 4. Discussion

The primary finding of this review is that the proportion of studies which failed to apply multilevel models to analyzing school-based data appears to be relatively high at 31.4%. In addition, even if multilevel models were applied, specification of levels of clustering effects was rarely described (data not shown). For instance, clustering effects of only the highest-level units of data hierarchy seem to have been taken into account, ignoring additional potential clustering effects of lower-levels units. Taken together, significance of findings based on $p$ values from those studies might have possibly been falsely declared especially when the $p$ values are close to 0.05. Furthermore, less than half of all studies reported power analysis, and approximately one-fifth of all studies, or equivalently about half of the studies that reported power analysis, took the ICC into consideration for power analysis. Therefore, it is likely that a majority of studies might have been underpowered implying that even if the study findings are significant, one cannot rule out the possibility of type I error.

The magnitudes of ICC for power analysis were mostly low, and rationales for such a hypothesized ICC are seldom clearly described. The dearth of rationale might have been due to the lack of information regarding ICC estimates, published or not, pertinent to their studies. This is reflected on the very low proportion of studies (8.3%) that reported ICCs estimated from their data analysis. To this end, the
reporting of the ICC estimated from data analysis would be critical for designing future studies with adequate sample sizes. Even if the ICC appears to be very small, it needs to be accounted for power analysis. For example, when the number of students is as small as 30 for each school, the number of required schools with ICC = 0.01 increases by 29% for the same power, compared to when ICC = 0, regardless of hypothesized effect sizes. Therefore, the impact of a small ICC on the sample size could be substantial.

Detailed design characteristics are often referred to protocol papers published earlier, and adequacy of power analysis was not evaluable in the text of outcome papers. Classification of such outcome papers as no report of power analysis might have underestimated the proportion of studies with power analysis because the protocol papers might have properly reported power analysis. However, we surmise that it is fairly rare for a school-based study to be conducted exactly as planned or designed in the protocol papers as school environments are dynamic and changes may alter aspects of the design or analysis plans including sample size determinations/power analysis, statistical methods, and outcome parameters. If outcomes papers do not clearly delineate these aspects, it may be unclear to know whether the power analysis in the earlier protocol paper would be appropriate for the applied statistical methods for the analysis of trial outcomes. We believe that this issue would be a research topic worthy to investigate. After all, the following key design elements should be reported in the outcome analysis papers: target power, significance level, hypothesized effect sizes and ICCs and their rationales, the number of clusters, the number of levels, anticipated attrition rates, and the planned sample size of the subjects. This description can enable readers to effectively and clearly evaluate whether the study has been analyzed as designed, not being forced to compare with detailed elements described in the protocol papers.

Many trials are excluded from evaluation due to the utilization of single schools. These studies are mostly college-based trials likely because multicollage trials may be difficult to conduct compared to the other types of schools. Nonetheless, it is not possible to take the clustering effect into account not only for power analysis but also for statistical analysis, nor is it possible to estimate ICCs from data analysis from single-school/cluster studies. Although it could bring up an issue as to what extent clustering effects should be considered in general, this limitation should be addressed in regard to limited generalizability or transportability of the findings from single-school trials. Identical or similar findings from replicate studies in other school settings would validate the findings.

Our findings also have implications for doctoral programs training future obesity researchers, particularly those conducting school-based interventions. The programs include but are not limited to health, clinical, and school psychology subspecialties. Researchers should ensure that the most rigorous and appropriate methodologies, including issues related to clustering addressed in this study, are included as part of the core curriculum. In this way, students learn early in their careers the practice of reporting such information. To this point, the American Psychological Association (APA) recently released two APA Publications and Communications Board Task Force reports addressing standards for reporting study results. Separate reports were made for quantitative studies [26], as well as qualitative, meta-analytic, and mixed-methods research [27].

Although our review is confined to school-based trials, the findings may be applicable to other cluster randomized trials using different settings and different types of interventions and treatments in other research areas [28–31]. Collectively, therefore, it would be ideal for reports of cluster randomized trials in general to adhere to the aforementioned 2012 updated CONSORT guideline [24] for both outcome and protocol papers. This guideline proposes all the design and analysis elements that should be reported in a standardized manner on manuscripts based on cluster randomized trials. If reports are standardized, it will be beneficial for researchers not only to plan or design school-based trials or other cluster randomized trials but also to more efficiently conduct systematic reviews and meta-analyses with greater statistical power and clearer transparencies.

4.1. Limitations. There are limitations that should be counted when interpreting results from this review. First, the search strategy was rather incomprehensive including only PubMed papers, and the keywords for search may be coarse. Therefore, studies that would have been eligible and added to our evaluations might not have been captured, and the scopes of potentially missing studies are unknown. As is the case for review papers in general, subjective ratings may not be completely avoided even if efforts are placed in minimizing misclassification errors. For instance, we did not evaluate whether potential confounding factors were appropriately controlled for in the data analysis. This evaluation might rely more on subjective judgments with substantial knowledge on the research topics under study and also might be difficult to reach a consensus. Lastly, again, the proportion of the studies with the reported power analysis might have been underestimated because studies that referred detailed power analysis to a protocol paper without a minimum level of description in the text were counted as papers with no report of power analysis.

5. Conclusions

In conclusion, the extent of the application of multilevel models to analyzing school-based trials appears to have so far been inadequate. Key elements such as the hypothesized ICC for power analysis or sample size determinations and the reported ICC estimated from data analyses of school-based trials are missing for a majority of studies. Future school-based trials should specifically address these issues at both design and analysis stages, preferably adhering to the extended CONSORT guideline to increase rigor and reproducibility of experimental settings and study findings. Clinical implications drawn based on the outcomes from school-based trials with rigorously well-performed design, conduct, and analysis would be the most useful to advance
knowledge for preventing and treating pediatric and adolescent obesity epidemic.

Data Availability

The data analyzed for the present analysis along with programming codes will be available upon reasonable request.

Disclosure

The opinions expressed are those of the authors and not necessarily of the NIH or any other organization.

Conflicts of Interest

Dr. David B. Allison has received personal payments or promises for the same from IKEA, Law Offices of Ronald A. Marron, Nestle, Paul Hastings LLP, and Tomasik Kotin LLC, and multiple NIH grants to teach, develop, apply, and evaluate statistical methods.

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Supplementary Materials

This material contains the list of 121 papers that we reviewed and evaluated for this paper. (Supplementary Materials)

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