# Experiment Critiques

Using the outline of Reporting Guidelines for Experimental Research below, summarize and critique each aspect of the experiment you were assigned to read.

Summarize means recording the relevant information from the experiment you read. You should note the page number of the article where you found the information so that you can easily find it again during class discussions.

Critique means discussing whether:

* the information for each item was easily found (e.g., were the hypotheses easy to identify? Was information about the experimental population and context provided?)
* each item was clearly explained (e.g., were the hypotheses easy to identify? Were hypotheses written clearly as a falsifiable statement? Could you replicate the experiment based on the description of treatments, procedures, analysis, etc?)
* each aspect of the experiment seems appropriate to answer the research question (e.g., does the population and context limit generalizability? Do the treatments seem to do what researchers intended [internal validity]?)
* whether you have concerns about the ethics of the experiment, including formal human subjects research principles, transparency/open science, or other ethical principles.

Note: You can skip the section of the original guidelines about creating a diagram of the research design (Section 5b) where the text is struck through.

*Source*

Gerber, Arceneaux, Boudreau, Dowling, Hillygus, Palfrey, Biggers and Hendry (2014). Reporting Guidelines for Experimental Research: A Report from the Experimental Research Section Standards Committee. *Journal of Experimental Political Science*, 1(1): 81-98. doi:10.1017/xps.2

1. **Hypotheses**
   1. State specific objectives or hypotheses.
   2. What question(s) was (were) the experiment designed to address?
   3. What are the specific hypotheses to be tested?
2. **Subjects and Context**
   1. Report eligibility and exclusion criteria for participants.
   2. Why was this subject pool selected? Who was eligible to participate in the study? What would result in the exclusion of a participant? Were any aspects of recruitment changed (such as the exclusion criteria) after recruitment began?
   3. Report procedures used to recruit and select participants.
   4. How were participants contacted for recruitment? Were incentives offered?
   5. If there is a survey: Identify the survey firm used and describe how they recruit respondents.
   6. Report recruitment dates defining the periods of recruitment and when the experiments were conducted.
   7. Also list dates of any repeated measurements as part of a follow-up.
   8. Describe settings and locations where the data were collected.
   9. In the field, lab, classroom, or some other specialized setting?
   10. Other relevant specifics of the population: e.g., large public university vs. small private university; geographic location; etc.
   11. If there is a survey: Provide response rate and how it was calculated.
3. **Allocation Method**
   1. Report details of the procedure used to generate the assignment sequence (e.g., randomization procedures).
   2. If random assignment used, report details of procedure (e.g., any restrictions, blocking).
   3. Note the unit of randomization (individuals, groups, households, etc.). Pay careful attention to report clustered random assignment if subjects were assigned at some level other than the individual subject.
   4. If random assignment used, to help detect errors such as problems in the procedure used for random assignment or failure to properly account for blocking, provide a table (in text or appendix) showing baseline means and standard deviations for demographic characteristics and other pretreatment measures (if collected) by experimental group.
   5. If blocking was used, and group assignment proportions were not equal across blocks, provide a table for each of the blocks. If there are too many blocks for this to be practical, combine blocks to present weighted averages of covariates using inverse probability weighting.
   6. Describe blinding.
   7. Were participants, those administering the interventions, and those assessing the outcomes unaware of condition assignments?
   8. If blinding took place, include a statement regarding how it was accomplished and how the success of blinding was evaluated.
4. **Treatments**
   1. Provide a detailed description of the interventions in each treatment condition, as well as a description of the control group.
   2. Descriptions should be sufficient to allow precise replication: Summary or paraphrasing of experimental instructions in the article text; verbatim instructions and/or other treatment materials provided in an appendix.
   3. State how and when manipulations or interventions were administered.
   4. Method of delivery: Pen-and-paper vs. computer or Internet vs. face-to-face communications vs. over the telephone.
   5. If computerized, the software should be described and cited. (If possible, programs should be included in an appendix so as to be available for purposes of replication.)
   6. For lab experiments (and other experiments, when relevant):
      1. Report the number of repetitions of the experimental task and the group rotation protocol. Report the ordering of treatments for within-subject designs. Any piggybacking of other protocols should be reported. Report any use of experienced subjects or subjects used in more than one session or treatment.
      2. Report time span: How long did each experiment last? How many sessions were subjects expected to attend? If there were multiple sessions, how much time passed between them?
      3. Report total number of sessions conducted and number of subjects used in each session.
      4. Report whether deception was used.
      5. Report treatment fidelity: Evidence on whether the treatment was delivered as intended.
      6. Report any instructional anomalies or inaccuracies.
      7. Were subjects given quizzes on the experimental instructions?
      8. Were there practice rounds? If so, how many and what were the results?
      9. Did subjects complete a post-experiment debriefing, interview, or questionnaire? If so, is there evidence that subjects understood the instructions and treatments?
      10. Did the experimental team observe aspects of the intervention?
      11. Provide descriptions of manipulation checks, if any.
      12. Were incentives given? If so, what were they and how were they administered?
5. **Results**
   1. **Outcome Measures and Covariates**
      1. Provide precise definitions of all primary and secondary measures and covariates.
      2. For indices, provide exact description of how they are formed. For survey items, provide exact question wording in an appendix. Provide a copy of the complete survey questionnaire (in an online appendix if it is long).
      3. Clearly state which of the outcomes and subgroup analyses were specified prior to the experiment and which were the result of exploratory analysis.
   2. **~~CONSORT Participant Flow Diagram~~**
      1. ~~Complete CONSORT Participant Flow Diagram~~
      2. ~~Note that the CONSORT flow chart entries include:~~
         1. ~~Number of subjects initially assessed for eligibility for the study.~~
         2. ~~Exclusions prior to random assignment and reasons for the exclusions.~~
         3. ~~Number of subjects initially assigned to each experimental group.~~
         4. ~~The proportion of each group that received its allocated intervention and the reasons why subjects did not receive the intended intervention.~~
         5. ~~The number of subjects in each group that dropped out or for other reasons do not have outcome data.~~
         6. ~~The number of subjects in each group that are included in the statistical analysis, and the reasons for any exclusions.~~
   3. **Statistical Analysis**
      1. Researchers will conduct statistical analysis and report their results in the manner they deem appropriate. We recommend that this reporting include the following:
      2. Report sample means and standard deviations for the outcome variables using intent-to-treat (ITT) analysis (means for the entire collection of subjects assigned to a group, whether the treatment is successfully delivered or not).
      3. If the experiment uses block randomization with unequal assignment rates, present ITT analysis by block or present overall means using inverse probability weighting.
      4. Note whether the level of analysis differs from level of randomization and estimate appropriate standard errors.
      5. If there is attrition, discuss reasons for attrition and examine whether attrition is related to pretreatment variables.
      6. Report other missing data (not outcome variables):
         1. Frequency or percentages of missing data by group.
         2. Methods for addressing missing data (e.g., listwise deletion, imputation methods).
         3. For each primary and secondary outcome and for each subgroup, provide summary of the number of cases deleted from each analysis and rationale for dropping the cases.
      7. For survey experiments: Describe in detail any weighting procedures that are used
6. **Other Information**
   1. Provide additional information about the experiment.
   2. Was the experiment reviewed and approved by an IRB?
   3. If the experimental protocol was registered, where and how can the filing be accessed?
   4. What was the source of funding? What was the role of the funders in the analysis of the experiment?
   5. Were there any restrictions or arrangements regarding what findings could be published? Are there any funding sources where conflict of interest might be an issue?
   6. If a replication data set is available, provide the URL