PRS Efficacy Criteria for Best-Evidence Risk Reduction (RR) Community-level Interventions (CLIs)

Intervention Description
• Clear description of key aspects of the intervention

Quality of Study Design
• Prospective study design
• Appropriate and concurrent control/comparison arm
• ≥ 4 communities per arm or appropriate power analysis indicating that a smaller number of communities was adequate (i.e., 2 or 3 communities per arm)
• Select similar communities (units) for assignment
  o To minimize selection bias before assignment regardless of assignment methods (randomization or not); use methods such as systematic, a priori approaches to choose intervention and control communities that are similar (e.g., matching or stratification on factors related to important/appropriate community characteristics)

Quality of Study Implementation and Analysis
• Sample individuals from assigned communities in acceptable ways (e.g., random, systematic) and use identical methods and eligibility criteria for selecting participants in each community, study arm, and data collection wave
  o If demographic differences are identified a priori, differential selection (e.g., over-sampling based on demographics) may be used to achieve equivalence between study arms on those factors
• Follow-up assessment ≥ 3 months post completion of entire time-specific CLI or post full implementation of on-going CLI with recall not referring to pre-intervention period
  o “Post full implementation of an on-going CLI” means after all components of the CLI have been started or put in place in communities
• If cohort, at least 70% retention rate at a single follow-up assessment for each study arm
  o If cohort chart review, ≥ 70% success rate in matching medical records
• Comparison between intervention arm and an appropriate comparison arm
• Analysis of communities (units) and analysis of individuals within the communities as originally assigned regardless of contamination or logistic/implementation issues
• Analysis of communities (units) regardless of community level of intervention exposure
• Analysis of individuals within the communities (units) regardless of individual level of intervention exposure
• Use of appropriate cluster-level analyses, e.g., adjusting for ICC
• Analysis must be based on post-intervention levels or among pre-post changes in measures
  o For pre-post changes used in analysis, measures must be identical, including identical recall period
• Analysis based on an a=.05 (or more stringent) and a 2-sided test
• Either no statistical differences in baseline levels of the outcome exist or baseline differences are controlled for in the analysis, regardless of allocation method (e.g., randomization, non-randomization)
  o No differences on baseline levels of the outcome means reporting no significant difference between groups on BL relevant outcomes or match/stratify/statistically adjust participant data by using propensity scores or relevant outcome covariates (regardless of assignment methods - RCT or non-RCT)

**Strength of Evidence**

**Demonstrated Significant Positive Intervention Effects**
- Positive and statistically significant ($p < .05$) intervention effect for $\geq 1$ relevant outcome measure
  o A positive intervention effect is defined as a greater reduction in HIV/STD incidence or risk behaviors or a greater increase in HIV protective behaviors in the intervention arm relative to the comparison arm.
  o A relevant outcome is defined as a behavior (e.g., abstinence, mutual monogamy, number of sex partners, consistent condom use with anal/vaginal sex, unprotected anal/vaginal sex, proportion of anal/vaginal sex acts protected, injection drug use, sharing or borrowing needles/works) that directly impacts HIV risk or a biologic measure indicating HIV or STD infection (i.e., HIV or STD incidence)
- Effect at the follow-up and based on the analyses that meet study implementation and analysis criteria

**No Demonstrated Significant Negative Intervention Effects**
- No negative and statistically significant ($p < .05$) intervention effect for any relevant outcome
  o A negative intervention effect is defined as a greater increase in HIV/STD incidence or risk behaviors or a greater decrease in HIV protective behaviors in the intervention arm relative to the comparison arm.
  o No other statistically significant harmful intervention effect
  o For an intervention with a replication evaluation, no significant negative intervention effects in the replication study

**Additional Limitations to Evaluate**
- No evidence that additional limitations resulted in a fatal flaw:
  o A fatal flaw has occurred when the overall evaluation of limitations resulted in considerable bias, thus substantially reducing the confidence of the findings
  o Examples of limitations to check for possible fatal flaw:
    - Group non-equivalence in baseline measures of important demographics or risk factors
    - Differential Retention (for cohort studies): (1) association between study arms and characteristics related to retention or attrition; OR (2) more than minimal rate of differential retention (> 10%)
- Differential Refusal: At baseline for cohort studies; by wave for serial cross-sectional studies: (1) association between study arms and characteristics related to refusal; OR (2) more than minimal rate of differential refusal rate (> 100)
- Intervention activities did not match with the intervention concepts or guiding theories intended to produce the desired outcomes
- Did not clearly describe issues related to generalizability
- Effects only found within a potentially biased subset analyses
- Substantial missing data (> 10% or missing data plus loss to attrition does not exceed acceptable limits for retention alone)
- Too many post hoc analyses (even with Bonferroni corrections)
- Pilot study or very small sample size per study arm (< 50)