SAFETY COUNTS
Good Evidence – Risk Reduction

INTERVENTION DESCRIPTION

Target Population
• Out-of-treatment active crack and injection drug users

Goal of Intervention
• Eliminate or reduce sex risk behaviors
• Eliminate or reduce drug risk behaviors

Brief Description
The Safety Counts intervention consists of a total of 9 sessions focusing on developing and implementing a personalized risk reduction plan. First, 2 individual standard pre- and post-test counseling sessions incorporate drug-focused prevention education to review basic HIV/AIDS information and provide optional HIV testing and counseling. Next, 2 interactive group workshop sessions, employing stages of change framework, are implemented with structured exercises involving 3-7 clients to help them develop a personal HIV risk reduction plan, consider potential barriers and solutions, identify sources of social support through group discussion, view role model videos, and complete 2 worksheet exercises to identify their own HIV risks and place themselves in on a stages-of-change continuum for each risk behavior. Then a one-on-one individual counseling session is conducted to refine the client’s personal risk reduction plan, strengthen commitment to personal goals, ensure availability of social support for risk reduction, and assess and arrange referral needs. One month after the client receives the individual counseling session, a minimum of two 15-20 minute field-based supportive follow-up outreach contacts are scheduled to reinforce progress toward risk reduction and encourage achievement and maintenance of personal risk reduction goals. Also, a minimum of 2 monthly social events, each lasting 2 hours, are provided, including lunch and planned HIV risk reduction activities, games, and skits for clients and their peer support buddies (15-25 clients and 10-15 guests) to provide support for HIV risk reduction, influence perceived social norms, and increase self-efficacy for reducing HIV risks. Lastly, food bank grocery bags and food coupons are made available to clients in storefront offices as a program incentive every other week.

Theoretical Basis
• Health Belief Model
• Theory of Protection Motivation
• Transtheoretical Stages of Change Model

Intervention Duration
• Nine sessions over a 4-6 month period
Intervention Setting
- Small businesses, neighborhood organizations, social agencies, on the street, and in other community settings

Deliverer
- Trained outreach specialists, a network of peer community volunteers, and a full-time coordinator supervising the peer network

Delivery Methods
- Counseling
- Exercise
- Games
- Goal setting/plan
- Group discussion
- Role play
- Video

INTERVENTION PACKAGE INFORMATION

In August 2013, the Centers for Disease Control and Prevention’s Division of HIV/AIDS Prevention (DHAP) announced that in accordance with its High Impact Prevention approach, DHAP will focus its behavioral intervention portfolio on interventions that are cost-effective, scalable and prioritize prevention for persons living with HIV and those persons at highest risk for acquiring HIV. Safety Counts will no longer be funded by DHAP for diffusion, adoption, and implementation.

Researchers: Dr. Scott Hershberger and Dr. Fen Rhodes have retired from the University of California at Long Beach. At this time there is no current contact information for this intervention.

EVALUATION STUDY AND RESULTS

The original evaluation was conducted in Long Beach, California, between January 1992 and December 1996. This was one of the 23 cooperative agreement studies under the NIDA Cooperative Agreement for AIDS Community-Based Outreach/Intervention Research Program.

Key Intervention Effect
- Reduced drug injection
- Reduced sharing works

Study Sample
The analytic study sample of 726 drug users is characterized by the following:
- 47% black or African American, 28% white, 20% Hispanic/Latino, 4% Native American, 1% Asian
- 67% male, 33% female
- Mean age of 39 years
- 59% completed high school education
Recruitment Settings
Street outreach in public areas and the community

Eligibility Criteria
Men and women were eligible if they were at least 18 years old, reported injecting drugs or using crack cocaine in the past 30 days, provided confirmation of recent drug use through urine based drug testing or track marks, and were not in drug treatment in past 30 days.

Assignment Method
The community from which participants were sampled was divided into two comparable geographic regions, each containing 3 zip codes. One region was randomly selected to receive the intervention initially, with one crossover midway through the study. A total of 1,362 eligible drug users were then assigned to 1 of 2 groups based on their “hangout” zip code: Safety Counts enhanced intervention (n = 687) or NIDA standard comparison (n = 675).

Comparison Group
The NIDA standard intervention was delivered in public areas and communities to individuals, or small groups or pairs of drug users for outreach. The intervention was delivered in two 20-30 minute sessions by an indigenous peer outreach worker and counselor, and included counseling, skills building, drug-focused prevention education as mandated by NIDA to review basic HIV/AIDS information, optional HIV testing, and referral to other services.

Relevant Outcomes Measured and Follow-up Time
- Sex behaviors during past 30 days include: having any sex, percentage of times used condoms, percentage of times always used condoms, having 2 or more sex partners, exchanged sex for drugs, and having sex with an IDU.
- Needle-related risk behaviors during past 30 days include: injecting any drugs, number of times injected, percentage of times did not use own works, and percentage of times used unclean needles.
- Sex and drug outcomes were measured at 5 to 9 months after baseline, which translates to 1 to 5 months after intervention.

Participant Retention
- Enhanced Safety Counts Intervention
  - 74% retained at 1-5 months after intervention
- Standard Intervention
  - 76% retained at 1-5 months after intervention

Significant Findings
- The participants in the Enhanced intervention were significantly less likely to report injecting drugs (p < 0.05) than those in the Standard at 1 to 5 months after intervention.
- Among injectors only, the percentage of times people did not use their own works was significantly lower in the Enhanced intervention compared to the Standard at 1 to 5 months after intervention (p < 0.05).
Considerations

- This intervention fails to meet the best-evidence criteria due to assigning groups of individuals to study conditions while analyzing at the individual level, a small number of participants being excluded from analyses after assignment, and a short follow-up time.
- There were three significant baseline demographic differences. The standard intervention group included more Hispanics (23% vs. 19%), fewer Asians (0.6% vs. 2%), and fewer married people (8% vs. 12%) than the Enhanced intervention group.
- Of the 687 participants assigned to the Enhanced intervention group, 462 (67%) did not receive all 9 sessions as allocated, whereas only 61 (9%) of the 675 participants assigned to the Standard intervention participants did not receive the full 2 sessions as allocated.
- Among those that completed the intervention as allocated, participants in the Enhanced group were significantly less likely to report having sex at follow-up compared to those in the standard group (p < .05). This finding does not satisfy good-evidence efficacy criteria due to a potentially biased restriction based on complete exposure.
- Among those that completed at least 7 out of 9 sessions, participants in the Enhanced group were significantly more likely to report an increase in condom use from baseline to follow-up as compared to those in the Standard group (p = .01). This finding does not satisfy good-evidence efficacy criteria due to a potentially biased restriction based on complete exposure.
- Among injectors that completed at least 7 out of 9 sessions, participants in the Enhanced group were significantly more likely to report decreases in high-risk drug behaviors from baseline to follow-up – stopped injecting drugs, p < .001, decreased number of days injected drugs, p = .001, decreased frequency of injecting drugs, p < .001 – as compared to those in the Standard group. These findings do not satisfy good-evidence efficacy criteria due to a potentially biased restriction based on complete exposure.
REFERENCES AND CONTACT INFORMATION


**Researchers: Dr. Scott Hershberger** and **Dr. Fen Rhodes** have retired from the University of California at Long Beach. **At this time there is no current contact information for this intervention.**