Interventions to address unequal gender and power relations and improve self-efficacy and empowerment for sexual and reproductive health decision-making for women living with HIV: A systematic review

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Abstract

Background

Many women living with HIV experience gendered power inequalities, particularly in their intimate relationships, that prevent them from achieving optimal sexual and reproductive health (SRH) and exercising their rights. We assessed the effectiveness of interventions to improve self-efficacy and empowerment of women living with HIV to make SRH decisions through a systematic review.

Methods and findings

We included peer-reviewed articles indexed in PubMed, PsycINFO, CINAHL, Embase, and Scopus published through January 3, 2017, presenting multi-arm or pre-post intervention evaluations measuring one of the following outcomes: (1) self-efficacy, empowerment, or measures of SRH decision-making ability, (2) SRH behaviors (e.g., condom use, contraceptive use), or (3) SRH outcomes (e.g., sexually transmitted infections [STIs]). Twenty-one studies evaluating 11 intervention approaches met the inclusion criteria. All were conducted in the United States or sub-Saharan Africa. Two high-quality randomized controlled trials (RCTs) showed significant decreases in incident gonorrhea and chlamydia. Sixteen studies measuring condom use generally found moderate increases associated with the intervention, including in higher-quality RCTs. Findings on contraceptive use, condom self-efficacy, and other empowerment measures (e.g., sexual communication, equitable relationship power) were mixed. Studies were limited by small sample sizes, high loss to follow-up, and high reported baseline condom use.
Conclusions

While more research is needed, the limited existing evidence suggests that these interventions may help support the SRH and rights of women living with HIV. This review particularly highlights the importance of these interventions for preventing STIs, which present a significant health burden for women living with HIV that is rarely addressed holistically. Empowerment-based interventions should be considered as part of a comprehensive package of STI and other SRH services for women living with HIV.

Introduction

An increasing body of evidence demonstrates the ways unequal levels of power between men and women in intimate relationships prevent women, including women living with HIV, from making decisions regarding their sexual and reproductive health (SRH) [1–5]. Gender refers to the set of roles, behaviors, and norms that are designated as appropriate for women and men by society [6]. Gender can be the cause, consequence, and/or mechanism of unequal or hierarchical power relations—that is, how power and control are distributed (unequally or hierarchically) in intimate relationships, within the household, in the community, and in wider societal institutions including all the way to the highest levels of political decision-making [6]. In this paper, we focus primarily on the distribution of power in intimate relationships between women and men and within the household. Frequently, unequal control over and access to economic resources, unequal relationship power, and limited ability to make sexual decisions (including whether, when, how often, and with whom to have sex; and negotiating condom use, contraceptive or other protective practices) make women vulnerable to SRH risks [7,8]. Gender inequalities and power imbalances restrict the ability of many women living with HIV to meet their SRH needs and exercise their rights [9].

One approach to address gender inequalities is implementing interventions that seek to empower women living with HIV. Empowerment has been defined as “the process of enhancing the capacity of individuals or groups to make choices and to transform those choices into desired actions and outcomes” [10]. Such interventions are designed to increase women’s self-efficacy, autonomy, or agency, and, hence, improve their sexual and reproductive decision-making and related health outcomes. However, although some interventions have been evaluated on an individual basis, the effectiveness of such interventions as a whole has not been systematically assessed through meta-analyses or systematic reviews.

We conducted a systematic review to examine the effectiveness of interventions that aim to address unequal gender power relations, empower women living with HIV, and increase their self-efficacy to make SRH decisions.

Methods

This systematic review was conducted to inform World Health Organization guidelines on the sexual and reproductive health and rights of women living with HIV, following PRISMA reporting guidelines [11]; the review protocol is available upon request [12].

Eligibility criteria

Studies were eligible for inclusion if they met the following criteria:
1. Examined one or more interventions designed to address unequal gender power relations, increase self-efficacy, and/or increase empowerment around safer sex and reproductive decision-making for women living with HIV,

2. Compared women living with HIV who received the intervention to those who did not through a pre/post or multi-arm design,

3. Measured at least one of the following outcomes: (a) Self-efficacy, empowerment, or other measure of ability to make own decisions around condom use, pregnancy termination, birth spacing, childbearing, and other aspects of SRH, (b) SRH behaviors (such as condom use, contraceptive use, disclosure of HIV serostatus to partner) or (c) SRH outcomes (such as STIs, pregnancy).

4. Published in a peer-reviewed journal prior to the search date.

We included studies among all populations of women living with HIV, including adolescents (10–19 years), young people (20–24 years), adults (25+ years), and women of any age who were members of key populations (including female sex workers, women who use drugs, women in prisons or other closed settings, and transgender women) [13]. Given our focus on SRH decision-making, we excluded studies with children under ten years of age. If a study evaluated an intervention for both men and women, or for both women living with HIV and HIV-negative women, it was included only if outcome data were disaggregated for women living with HIV. We did not include self-efficacy for coping with HIV status; self-efficacy for adherence to medications; or general measures of self-efficacy, self-esteem, agency, or wellness not directly linked with SRH behaviors and outcomes. Articles from all countries and written in all languages were eligible for inclusion.

Data sources
The following electronic databases were searched for articles through January 3, 2017: PubMed, CINAHL, Embase, PsycINFO, and Scopus. We developed search terms for HIV, women, study design, and SRH to identify articles in PubMed (S1 Appendix), then adapted the search for other databases. Secondary reference searching was conducted on all included articles.

Data analysis
Initial screening of titles and abstracts was conducted by JR and SM. Potentially relevant citations were then independently screened in duplicate by JR and SM and resolved through discussion with CK. Full-text articles were reviewed for final eligibility decisions.

JR and SM independently extracted data in duplicate using standardized forms. Differences in data extraction were resolved through discussion and consensus. The following information was gathered from each included study: objectives, location, population characteristics, intervention description, study design, sample size, follow-up periods, loss-to-follow-up, analytic approach, outcome measures, comparison groups, effect sizes, confidence intervals, significance levels, conclusions, and limitations. JR and CK assessed study rigor using the Evidence Project’s tool for evaluating multiple study designs in HIV behavioral intervention research [14], including assessment of comparison groups, random assignment and selection, follow-up rate, equivalency of comparison groups, and control for potential confounders.

Data were descriptively analyzed by coding categories and SRH outcomes. We did not meta-analyze due to differences in intervention design and outcome measurement across
studies. However, we grouped similar measures (e.g., condom use self-efficacy) across studies and summarized findings by outcome.

Results

Database searches produced a total of 3,351 hits; 2,087 citations remained after removing duplicates (Fig 1). After initial screening, 151 citations were reviewed by two authors in duplicate, of which 73 were excluded for not meeting the inclusion criteria (e.g., qualitative studies, studies without relevant outcomes, or studies without findings for women living with HIV). Seventy-eight articles were pulled for full-text review, and 57 were excluded. Ultimately, 21 studies were included in the review covering 11 specific intervention approaches (Table 1).

Study descriptions

Location. Thirteen studies were located in the United States (US) [15–27], while eight were adapted from US-based interventions to an African context, including four in South
Table 1. Descriptions of included studies.

| Author Year Location | Sexual and Reproductive Health and Rights Empowerment Intervention | Study Design | Intervention Provider | Theoretical Framework | Study Outcomes* |
|----------------------|-------------------------------------------------|--------------|-----------------------|-----------------------|-----------------|
| **SISTA Adaptations (WILLOW, PURSE)** | | | | | |
| Wingood, et al., 2004 | SISTA Adaptations (WiLLOW, PURSE) | *WILLOW*: ‘Women involved in life learning from other women’ | RCT, individual | Trained female health educator, co-facilitated by HIV-positive female peer educator | Social cognitive theory; Theory of gender and power; designed for women living with HIV |
| | Atlanta, GA; Birmingham, AL, USA | 4, 4-hour interactive group sessions implemented over consecutive weeks with 8–10 group participants. | N = 366 | Follow-Up Time: 12 months | Unprotected vaginal intercourse |
| | | Topics Covered: Gender pride; supportive social network use and maintenance; HIV transmission risk behaviors, communication and safe sex negotiation, condom use, managing abusive relationships | | | Proportion never used condoms |
| | | Primary Objective: Reduce unprotected vaginal sex | | | Incident STDs |
| | | | | | Condom Self-Efficacy |
| Saleh-Onoya et al., 2009 | SISTA Adaptations (WiLLOW, PURSE) | WiLLOW Adaptation | RCT, individual | Black, isiXhosa speaking, female health educator and a black isiXhosa speaking HIV-positive woman co-facilitator | Social cognitive theory; Theory of gender and power; designed for women living with HIV |
| | Western Cape, South Africa | 4, 4-hour group sessions implemented over consecutive weeks with 8–10 group participants | N = 120 | Follow-Up Time: 3 months | Self-efficacy for negotiating condom use |
| | | Topics Covered: sexual risk reduction and coping training (e.g., ethnic and gender pride, self-esteem, support networks, communication, HIV risk behaviors, etc.) | | | Self-efficacy for correct condom use |
| | | Primary Objective: Enhance coping skills and consistent condom use | | | Control in relationships |
| Klein et al., 2013 | SISTA Adaptations (WiLLOW, PURSE) | Multimedia WiLLOW | RCT, individual | Interactive computer modules with female African American narrator | Social cognitive theory; Theory of gender power, built from each piece of WiLLOW meetings |
| | Southern USA | 2, 1-hour interactive computer session separated into 2–8 minute activity modules | N = 175 | Follow-Up Time: 3 months | Condom Use |
| | | Topics covered: pride, values, goals, using social support, stress management, risk reduction, condom management, building healthy relationships, HIV re-infection, STIs, partner communication, disclosure, condom self-efficacy, computer use instructions | | | Partner sexual communication |
| | | Primary Objective: Increase protective sexual behaviors and psychosocial mediators associated with HIV risk reduction | | | Communication self-efficacy |
| Sarnquist et al., 2014 | SISTA Adaptations (WiLLOW, PURSE) | PURSE: ‘Peers Undertaking Reproductive and Sexual Health Education’ | Non-randomized trial | Nurses with enhanced FP training | Social learning theory, Theory of gender and power |
| | Chitungwiza, Zimbabwe | 3, 90-minute group sessions | N = 98 | Follow-Up Time: 3 months postpartum | Relationship power |
| | | Topics covered: sexual negotiation skills and empowerment, information about HIV, PMTCT, and FP, and communication skills related to sex and FP. | | | Control over condom use |
| | | Primary Objective: Achieve desired family size and spacing; maximize maternal and child health | | | Long-acting reversible contraception (LARC) use |

*Continued*
## Table 1. (Continued)

| Author Year Location | Sexual and Reproductive Health and Rights Empowerment Intervention | Study Design | Intervention Provider | Theoretical Framework | Study Outcomes* |
|-----------------------|---------------------------------------------------------------|--------------|-----------------------|-----------------------|-----------------|
| **SWP SMART/EST Women’s Project; NOW/NOW2; The Partner Project** |
| Jones et al., 2001 | Miami, FL; Newark, NJ; New York, NY, USA | NOW: ‘New Opportunities for Women’ | Non-randomized trial with matched controls | Psychologist | Use of N-9 spermicides |
| | | • 3, 120-minute sessions over 3 months | · N = 178 |
| | | • Topics Covered: HIV/STD transmission, hierarchical counseling, skill training, reactions to barriers, cognitive reframing, and sexual negotiation | · Follow-Up Time: 9 months |
| | | • Primary Objective: Increase sexual barrier use | |
| | Jones et al., 2005 | Lusaka, Zambia | The Partner Project (NOW Adaptation) | RCT, individual | Theory of reasoned action/planned behavior |
| | | | • 4 group intervention sessions; male partners attended 1 or 4 separate sessions | · N = 332 (180 women living with HIV) |
| | | | • Topics covered: HIV/sexually transmitted disease prevention and transmission, reproductive choice and mother to child transmission, communication, conflict resolution, sexual negotiation | · Follow-Up Time: 12 months |
| | | | • Primary Objective: Reduce sexual risk behavior | |
| | Jones et al., 2006 | Lusaka, Zambia | NOW2 (NOW Adaptation) | RCT, individual | Theory of reasoned action/planned behavior |
| | | | • 2-hour group sessions limited to 10 women | · N = 240 |
| | | | • Topics covered: (1) HIV/STDs, safer sex, barrier use, reproductive choice, HIV re-infection, transmission and infection with other STDs and hierarchical methods of sexual barrier use (2) Vaginal lubricants, gels and suppositories | · Follow-Up Time: 12 months |
| | | | • Primary Objective: Increase sexual barrier use | |
| | Jones et al., 2007 | Miami, FL, USA | NOW ‘New Opportunities for Women’ | Randomized trial without control (randomized to individual or group sessions) | Facilitators were gender matched RNs, LPNs and health care staff trained in the administration of each condition |
| | | | • 3, 120-minute sessions over 3 months limited to 10 participants | · N = 187 |
| | | | • Topics covered: sexual barrier products, sexual risk reduction strategies, sexual negotiation | · Follow-Up Time: 12 months |
| | | | • Primary Objective: Increase sexual barrier use | |
| | Weiss et al., 2011 | | | |

(Continued)
| Author Year Location | Sexual and Reproductive Health and Rights Empowerment Intervention | Study Design | Intervention Provider | Theoretical Framework | Study Outcomes* |
|----------------------|--------------------------------------------------------------------|--------------|-----------------------|----------------------|-----------------|
| Miami, FL; Newark, NJ; New York, NY, USA | • SWP I and II ‘SMART/EST Women’s Program’ + Group Healthy Living Component • 10 weekly 2-hour group cognitive—behavioral stress management/exppressive—supportive therapy framework (CBSM+) • 6 additional 2-hour group healthy living sessions • Topics covered: medication adherence, nutrition, safer sex, substance abuse reduction, and physical activity. • Primary Objective: Optimize health status of poor women of color living with HIV | RCT, individual • N = 933 • Follow-Up Time: 24 months | Psychologist | Cognitive behavioral stress management (CBSM) plus expressive supportive therapy framework (CBSM+) | • Unprotected sex • Vaginal sexual barriers |
| Jones et al., 2013 | Miami, FL; Newark, NJ; New York, NY, USA | • SWP ‘SMART/EST Women’s Program’ Community Health Center Adaptation • 10 weekly 2-hour group cognitive—behavioral stress management/exppressive—supportive therapy framework (CBSM+) • 6 additional 2-hour group healthy living sessions • Topics covered: medication adherence, nutrition, physical activity, sexual risk behavior, and alcohol and drug use • Primary Objective: Optimize health status of women living with HIV in a community health setting | Non-randomized trial • N = 428 • Follow-Up Time: 12 months | Health-care providers, counselors, social workers, and health educators | Cognitive behavioral stress management (CBSM) plus expressive supportive therapy framework (CBSM+), Glasgow’s RE-AIM model | • Number of sexual partners |
| M2M ‘Mothers 2 Mothers’; Mamekaya | Peri-urban Cape Town, South Africa | • Mamekaya, based on M2M ‘Mothers 2 Mothers’ • 8 session, small groups of pregnant women • Topics Covered: Healthy Living-staying in care, dealing with symptoms, HIV, ARVs, family planning, condoms; Feeling Happy & Strong-disclosure, stigma, support, hope, negative emotions, domestic violence, substance abuse; Partnering & Preventing Transmission: infant feeding, partner testing, safer sex; Parenting: feeding, immunizations, infant testing, custody, attachment; in all sessions: music, meditation, active learning • Primary Objective: PMTCT and maternal well-being | Non-randomized trial, group • N = 160 • Follow-Up Time: 6 months post-delivery | M2M mentor mothers (women living with HIV) trained in CBI | Cognitive behavioral interventions, empowerment and support model | • Partner testing • Abstinence/condom use |
| Richter et al., 2014 | | | | | | (Continued) |
| Author Year Location | Sexual and Reproductive Health and Rights Empowerment Intervention | Study Design | Intervention Provider | Theoretical Framework | Study Outcomes* |
|----------------------|-------------------------------------------------|--------------|-----------------------|-----------------------|-----------------|
| KwaZulu-Natal, South Africa | **Mothers2Mothers Adaptation**<br>• 8 individual mentor sessions: 4 antenatal, 4 postnatal<br>• Topics covered: destigmatizing HIV, PMTCT tasks, exclusive feeding, abstaining from traditional medicines, healthy daily routines, obtaining a child grant, maintaining strong social network, couples’ HIV testing, disclosure, condom use<br>• Primary Objective: Maternal and infant well-being | RCT, group<br>• N = 1,200<br>• Follow-Up Time: 1.5 months post-birth | Peer mentors | Empowerment and support model | *Asking partner to test for HIV* |
| Holstad et al., 2011 | **KHARMA ‘Keeping Healthy and Active with Risk Reduction and Medication Adherence’**<br>• KHARMA ‘Keeping Healthy and Active with Risk Reduction and Medication Adherence’<br>• 8 group sessions<br>• Topics covered: ART adherence, risk behavior, HIV status disclosure<br>• Primary Objective: Promotion of adherence to antiretroviral medications and risk reduction behaviors | RCT, individual<br>• N = 203<br>• Follow-Up Time: 9 months | Trained nurses | Motivational interviewing theory | Abstinence<br>• Use of Protection |
| Holstad et al., 2012 | **KHARMA Adaptation**<br>• 8 group sessions<br>• Topics covered: ART adherence, self-efficacy for condom skills and knowledge, condom negotiation, HIV status disclosure<br>• Primary Objective: Promotion of adherence to antiretroviral medications and risk reduction behaviors | RCT, individual<br>• N = 60<br>• Follow-Up Time: 6 months | Trained nurses | Motivational interviewing theory; Social cognitive theory | Number of sexual partners<br>• Use of condoms/ protection<br>• Drug/alcohol use prior to sex |
| Marhefka et al., 2014 | **HR ‘Healthy Relationships’**<br>• HR-VG ‘Healthy Relationships—Videoconferencing Groups’ (HR Adaptation)<br>• 6, 2-hour videoconference sessions<br>• Topics covered: HIV status, disclosure decision-making and safer sexual behaviors<br>• Primary Objective: Reducing sexual risk behavior | RCT, individual<br>• N = 71<br>• Follow-Up Time: 6 months | 2 women living with HIV (1 social worker, 1 community member) | Social cognitive theory | Unprotected sex |
| Echenique et al., 2013 | **Project ROADMAP ‘Reeducating Older Adults in Maintaining AIDS Prevention’** | | | | |

(Continued)
Table 1. (Continued)

| Author Year Location | Sexual and Reproductive Health and Rights Empowerment Intervention | Study Design | Intervention Provider | Theoretical Framework | Study Outcomes* |
|----------------------|---------------------------------------------------------------|--------------|-----------------------|-----------------------|-----------------|
| Miami, FL USA         | • ROADMAP ‘Reeducating Older Adults in Maintaining AIDS Prevention’  
|                      | • 4 weekly psycho-educational group sessions for older women, 2-hours each  
|                      | • Topics covered: HIV, harm reduction, effects of HIV on sexual behaviors, assertive communication with partners, condom negotiation, de-escalating negative partner reactions, review of lessons learned, self-reward for maintaining safer behavior  
|                      | • Primary Objective: Reduce high risk sexual behavior  | • RCT, individual  
|                      | • N = 300  
|                      | • Follow-Up Time: 6 months  | Peer educators  | Information-motivation-behavioral skills (IMB model) of AIDS risk behavior change; principles of self-efficacy theory  | • Condom use  |
| FDIP ‘Women and Infants Demonstration Project’ | Fogarty et al., 2001  | • WDIP ‘Women and Infants Demonstration Project’  
|                      | • Unlimited individual sessions over 6 month period  
|                      | • Topics covered: condom and contraceptive use, condom negotiation  
|                      | • Primary Objective: condom and contraceptive use  | • RCT, individual  
|                      | • N = 322  
|                      | • Follow-Up Time: 18 months  | Trained peer mentors  | Stages of change theory  | • Condom use self-efficacy  
|                      |                                                             |                                                             | • Condom use  
|                      |                                                             |                                                             | • Contraceptive use  |
| Protect and Respect   | Teti et al., 2010  | • Protect and Respect  
|                      | • 5 consecutive, weekly, 1.5 hour group intervention sessions and peer-led support groups  
|                      | • Topics covered: sexual risk reduction education and skill-building; women’s challenges and opportunities; HIV/AIDS and STI facts; male and female condom use and condom negotiation; triggers to unsafe sex; HIV status disclosure; problem solving; healthy relationships; social support; and goal setting.  
|                      | • Primary Objective: increase HIV status disclosure and condom use  | • RCT, individual  
|                      | • N = 184  
|                      | • Follow-Up Time: 18 months  | Health care professionals, health educators, and peer educators  | Transtheoretical model of the stages of change; Modified AIDS risk reduction model; Theory of gender and power; formative research  | • Condom use  |
| WHC ‘Women’s Health CoOp’ | Wechsberg et al., 2010  | • WHC-Pretoria ‘Women’s Health CoOp’ (WHC Adaptation)  
|                      | • 2 individual 1-hour sessions held within a 2-week period  
|                      | • Topics covered: substance abuse, HIV/STIs, HIV risk, behavioral skills training with condoms, violence prevention, sexual negotiation and communication  
|                      | • Primary Objective: reduce sexual risk, substance use, and victimization among at-risk and underserved women  | • RCT, individual  
|                      | • N = 214  
|                      | • Follow-Up Time: 6 months  | Trained interventionist  | Gender and empowerment theories  | • Condom use  |
| ESHI ‘Enhanced Sexual Health Intervention’ | Wyatt et al., 2004  |                                                             |                                                             |                                                             |
Africa [28–31], two in Zambia [32,33], one in Zimbabwe [34], and one in Nigeria [35]. The US-based studies were largely implemented in urban areas. Two studies did not specify the exact study location, but were located in “a southern state with a high HIV prevalence” and “a large southeastern metropolitan city” respectively [16, 20].

**Population characteristics.** All studies included women living with HIV, per our inclusion criteria. Several studies focused on women from vulnerable or key populations, such as women with high rates of alcohol and other drug use [19,31], female sex workers [31], pregnant women [28,29,34], older adults [15], young women [27], and women with histories of child sexual abuse [26]. The US-based studies included primarily African-American and Hispanic women [15–26]. Across studies, ages ranged from 16–70 years old.

**Study design.** Tables 1 and 2 present information on study design and quality assessment. Sixteen studies were randomized controlled trials (RCTs) with randomization at either the individual or group (facility/community) level [15–17,21–27,29–33,35], while five studies employed other study designs, including non-randomized trials and a randomized trial with no control (participants randomized to group or individual intervention) [18–20,28,34]. Sample sizes at baseline ranged from 43 to 1,200; several of the smaller studies were described as feasibility or pilot studies. Follow-up time ranged from 3–24 months. Ten studies had follow-up rates of 75% or more.

**Theoretical bases.** All programs had an underlying theoretical basis. Theories used included social learning theory/social cognitive theory [15,21,22,25,30,34,35], the theory of gender and power [21,23,25,27,30,31,34], the empowerment and support model [28,29], the theory of reasoned action and theory of planned behavior [19,32,33], stages of change theory [16,23], and the AIDS risk reduction model [23].

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Table 1. (Continued)

| Author Year Location | Sexual and Reproductive Health and Rights Empowerment Intervention | Study Design | Intervention Provider | Theoretical Framework | Study Outcomes* |
|----------------------|---------------------------------------------------------------|--------------|-----------------------|-----------------------|---------------|
| **ESHI**  Enhanced Sexual Health Intervention  
Los Angeles, CA, USA  
- 11 weekly 2.5-hour psychoeducational sessions  
- Topics covered: HIV risk behaviors, interpersonal and health behaviors, and psychological symptoms  
- Primary Objective: reduce sexual risks and increase HIV medication adherence for HIV-positive women with child sexual abuse (CSA) histories | RCT, individual  
N = 147  
Follow-Up Time: 6 months | Trained group facilitator and peer mentor living with HIV with a history of CSA | Cognitive-behavioral approaches to risk reduction and cultural- and gender-specific concepts | Condom use |
| **EVOLUTION** Young Women Taking Charge and Growing Stronger  
Baltimore, MD; Chicago, IL; Tampa, FL, USA  
- 9 (7 group, 2 individual) weekly 2–3 hour sessions with 6–8 women per group  
- Topics covered: HIV risk reduction education and sexual negotiation skills, forgiveness, emotional regulation, communication, and relationships  
- Primary Objective: Decrease sexual risk; empower young women living with HIV through knowledge and skills | RCT, individual  
N = 43  
Follow-Up Time: 3 months | Trained group facilitator | Theory of gender and power | Sexual activity and sexual risk questionnaire  
Self-efficacy for limiting HIV risk behavior  
Self-efficacy for sexual discussion  
Condom use self-efficacy  
Sexual beliefs |

*Only outcomes relevant to self-efficacy and empowerment around sexual and reproductive health are included.

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Table 2. Quality assessment of included studies.

| Author, Year | Cohort | Control or comparison group | Pre/post intervention data | Random assignment of participants to the intervention | Random selection of participants for assessment | Follow-up rate of 75% or more | Comparison groups equivalent socio-demographics | Comparison groups equivalent at baseline on outcome measure | Control for potential confounders |
|--------------|--------|------------------------------|-----------------------------|---------------------------------------------------|---------------------------------|------------------------|---------------------------------|---------------------------------|-------------------------------|
| **SISTA Adaptations (Willow, PURSE)** | | | | | | | | | |
| Wingood et al., 2004 | Yes | Yes | Yes | Yes | No | Yes | Yes | Yes | Yes |
| Saleh-Onoya et al., 2009 | Yes | Yes | Yes | Yes | No | Yes | Yes | No | Yes |
| Klein et al., 2013 | Yes | Yes | Yes | Yes | No | Yes | Yes | Yes | Yes |
| Samquist et al., 2014 | Yes | Yes | Yes | No | No | Yes | Yes | Yes | Unclear |
| **SWP ‘SMART/EST Women’s Project’; NOW/NOW2; The Partner Project** | | | | | | | | | |
| Jones et al., 2001 | Yes | Yes | Yes | No | No | No | No | Yes | No |
| Jones et al., 2005 | Yes | Yes | Yes | Yes | No | Unclear | Unclear | Unclear | No |
| Jones et al., 2006 | Yes | Yes | Yes | Yes | No | No | No | Yes | No |
| Jones et al., 2007 | Yes | Yes | Yes | Yes | No | Unclear | Unclear | Unclear | No |
| Weiss et al., 2011 | Yes | Yes | Yes | Yes | No | Unclear | Unclear | Unclear | No |
| Jones et al., 2013 | Yes | Yes | Yes | No | Unclear | No | Yes | No | No |
| **M2M ‘Mothers 2 Mothers’; Mamekhaya** | | | | | | | | | |
| Futterman et al., 2010 | Yes | Yes | No | No | No | No | No | Unclear | Yes |
| Richter et al., 2014 | Yes | Yes | Yes | Yes | No | No | Yes | Yes | Unclear |
| **KHARMA ‘Keeping Healthy and Active with Risk Reduction and Medication Adherence’** | | | | | | | | | |
| Holstad et al., 2011 | Yes | Yes | Yes | Yes | No | Yes | No | Unclear | Yes |
| Holstad et al., 2012 | Yes | Yes | No | Yes | No | Yes | No | Unclear | Unclear |
| **HR ‘Healthy Relationships’** | | | | | | | | | |
| Marhefka et al., 2014 | Yes | Yes | Yes | Yes | No | Yes | Yes | Yes | Yes |
| **Project ROADMAP ‘Reeducating Older Adults in Maintaining AIDS Prevention’** | | | | | | | | | |
| Echenique et al., 2013 | Yes | Yes | Yes | Yes | No | No | Yes | Yes | No |
| **WDIP ‘Women and Infants Demonstration Project’** | | | | | | | | | |
| Fogarty et al., 2001 | Yes | Yes | Yes | Yes | No | No | Unclear | Unclear | Unclear |
| **Protect and Respect** | | | | | | | | | |
| Teti et al., 2010 | Yes | Yes | Yes | Yes | No | No | Yes | Yes | Yes |
| **WHC ‘Women’s Health Coop’** | | | | | | | | | |
| Wechsberg et al., 2010 | Yes | Yes | Yes | Yes | No | No | Unclear | Unclear | Yes |

(Continued)
Intervention descriptions. The 21 included studies covered 11 specific interventions (Table 1). Some interventions included multiple adaptations to different geographic context. In one instance, an in-person intervention was later adapted for multimedia [21]. Several were considered effective behavioral interventions by the U.S. Centers for Disease Control and Prevention.

Interventions were generally delivered in small group or one-on-one sessions. Several interventions incorporated cognitive-behavioral components, including cognitive-behavioral stress management/expressive-supportive therapy and cognitive-behavioral skill training [18,24,26,28]. Motivational interviewing was also common [15,17,35,36].

Study outcomes

Table 3 presents study outcomes. Two studies measured STI incidence [25,30]. Eighteen of the 21 studies measured sexual and reproductive health behaviors: 16 measured condom use [15–17,21–28,30–33,35] while two measured contraceptive use [16,34]. Six studies measured self-efficacy and psychosocial outcomes [16,21,25,27,30,34]. No studies measured reproductive health decision-making around pregnancy termination, birth spacing, or childbearing.

Sexually transmitted infections. Two studies measured STI incidence: the original WiLLow intervention in the southern USA and its South African adaptation. Both were high-quality RCTs, although the South African adaptation had a shorter follow-up time (3 vs. 12 months) and smaller sample size (102 vs. 321 participants) [25,30]. Both studies showed significant decreases in STI incidence. The original WiLLow intervention found a significant reduction in incidence of bacterial STIs (Chlamydia trachomatis and gonorrhea) over 12-month follow-up in intervention versus control participants (OR = 0.20, 95% CI = 0.10–0.60). However, there was no significant change in Trichomonas vaginalis [25]. In the South African adaptation, the intervention group similarly showed a significant reduction in incidence of Chlamydia trachomatis (OR = 0.21, 95% CI = 0.07–0.59) and gonorrhea (OR = 0.10, 95% CI = 0.02–0.49) compared to the control group. The South African adaptation further showed a significant decrease in incidence of Trichomonas vaginalis (OR = 0.06, CI = 0.01–0.46), but no difference in incidence of bacterial vaginosis [30].

Condom use. Sixteen studies (11 interventions) measured condom use [15–17,21–28,30–33,35]; however, studies used a wide range of measures, precluding meta-analysis. These studies (12 RCTs and four other designs) showed mixed results. Although most studies found significant increases in condom use, others found no change and increases were often moderate, often affected by high background rates of condom use.
Table 3. Sexual and reproductive health findings from included studies.

| Author Year | Study Findings |
|-------------|----------------|
| **SISTA Adaptations (WILLOW, PURSE)** | |
| Wingood et al., 2004 | • Condom use self-efficacy: 8.1 (95% CI 1.1, 15.0), p = 0.001  
  o % relative change comparing intervention to control: 8.1  
  (95% CI 1.1, 15.0), p = 0.001  
  o Adjusted mean difference: 1.0 (95% CI -28.0, -13.4), p = 0.022  
  o Adjusted mean difference: -0.7 (95% CI -1.8, -0.4)  
  • Number of acts of unprotected vaginal sex, past 30 days:  
  o % relative change comparing intervention to control: -28.0  
  (95% CI -69.3, -13.4), p = 0.008  
  o Proportion never used condoms, past 30 days:  
  o OR = 0.3 (95% CI 0.1, 0.7), p = 0.008  
  • Incident bacterial STD (chlamydia or gonorrhea):  
  o OR = 0.2 (95% CI 0.1, 0.6), p = 0.006  
  • Incident Trichomonas infection:  
  o No differences observed, no data reported |
| Saleh-Onoya et al., 2009 | • Condom use self-efficacy:  
  o F = 1.65, p = 0.20  
  • Self-efficacy for negotiating condom use:  
  o F = 0.47, p = 0.50  
  • Relationship power:  
  o F = 0.77, p = 0.38  
  • Condom use at last sex:  
  o OR = 0.48 (95% CI 0.09, 2.54), p = 0.39  
  • Number of unprotected vaginal sex, past 30 days:  
  o % relative change comparing intervention to control: -28.0  
  (95% CI -69.3, -13.4), p = 0.022  
  o Adjusted mean difference: -0.7 (95% CI -1.8, -0.4)  
  • Proportion never used condoms, past 30 days:  
  o OR = 0.3 (95% CI 0.1, 0.7), p = 0.008  
  • Incident bacterial STD (chlamydia or gonorrhea):  
  o OR = 0.2 (95% CI 0.1, 0.6), p = 0.006  
  • Incident Trichomonas infection:  
  o No differences observed, no data reported |
| Klein et al., 2013 | • Sexual communication self-efficacy:  
  o % relative change comparing intervention to control: 9.70  
  (95% CI 2.08, 21.77), p = 0.004  
  o Adjusted mean difference: 3.40 (95% CI 1.12, 5.65)  
  • Condom-protected vaginal and anal sex acts, past 30 days:  
  o % relative change comparing intervention to control: 45.21  
  (95% CI 17.67, 71.36), p = 0.002  
  o Adjusted mean difference: 0.33 (95% CI 0.13, 0.52)  
  • 100% condom use:  
  o OR = 9.67 (95% CI 1.25, 74.97), p = 0.30  
  • Number of unprotected vaginal and anal sex acts, past 30 days:  
  o % relative change comparing intervention to control: -133.67  
  (95% CI -190.20, -41.71), p = 0.002  
  o Adjusted mean difference: -3.41 (95% CI -5.54, -1.29)  
  • Disclosure of HIV serostatus, woman to partner:  
  o Intervention: 98.4%, Control: 55.2%, p = 0.04  |
| Sarnquist et al., 2014 | • Relationship power:  
  o Intervention: 2.5%, Control: 1.0%, p = 0.01  
  • Control over condom use:  
  o Intervention: 67.2%, Control: 34.4%, p = 0.002  
  • Use of long-acting reversible contraception:  
  o Intervention: 87.1%, Control: 81.8%, p = 0.34  
  • Use of N-9 spermicides:  
  o Intervention: 83%, Control: 9%, p<0.05  
  • Male condom use, 6 months after baseline:  
  o Group vs individual intervention: F = 13.5, p<0.001  
  • Male condom use, 12 months after baseline:  
  o Group vs individual intervention: F = 0.24, p = 0.62  
  • Number of sex partners:  
  o OR = 0.6 (95% CI 0.4–0.9)  |
| **SWP ’SMART/EST Women’s Project’; NOW/NOW2; The Partner Project** | |
| Jones et al., 2001 | • Use of N-9 spermicides:  
  o Intervention: 83%, Control: 9%, p<0.05  
  • Protected sex, 6 months after baseline:  
  o X = 4.90, t(1,70) = -.67, p = 0.001  |
| Jones et al., 2005 | • Protected sex, 12 months after baseline:  
  o X = 4.83, t(1,30) = -3.20, p = 0.003  
  • Male condom use, 6 months after baseline:  
  o Group vs individual intervention: F = 13.5, p<0.001  
  • Male condom use, 12 months after baseline:  
  o Group vs individual intervention: F = 2.4, p = 0.12  |
| Jones et al., 2006 | • Sexual barrier use, 6 months after baseline:  
  o Group vs individual intervention: F = 0.24, p = 0.62  |
| Jones et al., 2007 | • Sexual barrier use, 12 months after baseline:  
  o Group vs individual intervention: F = 0.5, p = 0.05  |
| Weiss et al., 2011 | • Unprotected sex:  
  o Decreased OR from 0.16 to 0.095, F = 0.04, p = 0.038  |
| Jones et al., 2013 | • Number of sex partners:  
  o OR = 0.6 (95% CI 0.4–0.9)  |
| **M2M ‘Mothers 2 Mothers’; Mamekhaya** | |
| Futterman et al., 2010 | • Abstinent or always uses condom:  
  o Coefficient: 0.24, SE: 1.44, p>0.05  |

(Continued)
Of the three SISTA adaptation RCTs that measured condom use, two showed significant increases [21,25], while the South African adaptation did not [30]. Most other high-quality RCTs also found significant positive impacts on condom use by various measurements [17,24,26,31–33,35]. WDIP found progress (through stages of change) in condom use with
main partner [16]. Three studies with high loss to follow-up rates (30–44% retention at follow-up) found mixed results on condom use [15,23,28].

**Contraceptive use.** Two studies measured contraceptive use. WDIP, an RCT, found that intervention participants were more likely to show progress (OR = 2.07, \( p = 0.08 \)) and significantly less likely to relapse (OR = 0.43, \( p = 0.03 \)) in contraceptive use compared to the comparison group [16]. PURSE, a non-randomized trial with 98 participants and high rates of follow-up, found that uptake of long-acting reversible contraception increased in both intervention and control groups three months after delivery, but there was no significant difference across groups (I: 87%, C: 81.8%, \( p = 0.34 \)). The authors suggested this was due to both groups having access to nurses with training in enhanced family planning [34].

**Self-efficacy and psychosocial measures.** Four RCTs (the original WiLLOW, its South African adaptation, WDIP, and EVOLUTION) and one non-randomized trial (PURSE) measured condom use self-efficacy. The original WiLLOW program found that intervention participants had higher condom use self-efficacy over 12 months of follow-up (13.6 vs. 12.6; \( p = 0.001 \)) [25]. PURSE also found significant increases in self-reported control over condom use (67.2% vs. 34.4%, \( p = 0.002 \)) [34], and WDIP intervention participants showed higher self-efficacy for condom use with a main partner than control participants (OR = 2.01, \( p = 0.01 \)) [16]. However, neither the small EVOLUTION pilot study nor the South African WiLLOW adaptation found a significant difference between intervention and control groups in condom use self-efficacy [30].

Other psychosocial outcomes also showed mixed results. The multimedia WiLLOW adaptation reported improvement in sexual communication self-efficacy (mean difference = 3.40, \( p = 0.004 \)) [21], while EVOLUTION found no significant impacts on self-efficacy for sexual discussion or self-efficacy for limiting HIV risk behavior [27]. PURSE found significant increases in relationship power (2.5 vs. 2.1, \( p = 0.01 \)) [34], whereas the South African WiLLOW adaptation found no significant results for relationship control or condom negotiation [30]. Finally, PURSE intervention participants were more likely to report disclosing their HIV status to a partner (98.4% vs. 87.5%, \( p = 0.04 \)) and vice versa (75.8% vs. 55.2%, \( p = 0.04 \)) [34].

**Discussion**

All women living with HIV must be supported in their voluntary choices around sexual relationships and be given information and resources to engage in safe, enjoyable sexual experiences, or to not engage in sex based on their personal preference, with counselling and support tailored to their decision-making, desires and needs. Supporting women living with HIV in all their diversity to achieve their sexual and reproductive health and rights in all epidemic contexts requires overcoming major barriers to service uptake such as social exclusion and marginalization, criminalization, stigma, and gender inequality [37]. Addressing unequal gender and power relations and empowering women living with HIV may be one part of a comprehensive approach to achieve these goals.

This systematic review highlights the potential for increasing condom use and reducing incident STIs through empowerment interventions for women living with HIV. STIs continue to be an important public health issue that can facilitate sexual transmission of HIV and trigger some cancers. As stated in the WHO Global Health Sector Strategy on Sexually Transmitted Infections, 2016–2021, “the burden of morbidity and mortality worldwide resulting from sexually transmitted pathogens compromises quality of life, as well as sexual and reproductive health” [37]. Women living with HIV have high rates of STI co-infection, with a mean STI prevalence of 15.8% (standard deviation: 9.9) across studies in a recent global systematic review [38]. Although STI screening and treatment are a recommended part of the package of
care for people living with HIV by the WHO [39,40] and PEPFAR [41], a comprehensive, rights-based approach to addressing STIs and other SRH issues is needed to facilitate STI prevention as well as treatment for women living with HIV.

Findings from our review were more mixed, however, for other outcomes, including contraceptive use, self-efficacy, and psychosocial measures. While these interventions hold promise, further work is needed to determine which components of interventions make them successful, for which populations, and on which outcomes.

Conclusions from this review are limited by the nature of the evidence base. The range of outcomes measured by the included studies was narrow, with the majority measuring condom use. Only a few studies measured other SRH outcomes, or more proximal outcomes such as empowerment and self-efficacy. Consequently, it is difficult to assess the impact of the interventions on women’s self-efficacy or empowerment, and to understand the association between empowerment and SRH outcomes. Not measuring other outcomes limits the evidence for pathways to improved health for women living with HIV and their partners. Additionally, studies used a wide range of measures for condom use that affected our ability to compare across interventions and precluded us from conducting meta-analysis. Condom use reported in these studies was affected by high rates of initial reported use, creating a ceiling for measuring intervention impact. Many measures were also self-reported, introducing the possibility of recall and social desirability bias. Finally, the included studies were of mixed quality, with many limited by small sample size and low follow-up rates. The evidence base is further limited in geographic and population scope. Many important populations of women living with HIV, such as transgender women, were not included in any studies. Most included studies were conducted in the USA or were adaptations of interventions originally implemented there. Nevertheless, some interventions were determined to be effective when adapted to multiple contexts and feasible across settings. Finally, we did not include unpublished (“grey”) literature or qualitative studies in our inclusion criteria; these studies may have provided additional insights into the effectiveness and outcomes of interventions.

Although this review focused on interventions with women, interventions with men that seek to address unequal gender and power relations are also essential to empower women in their SRH decisions. Recent evidence suggests that gender-transformative interventions to engaging men in HIV [42] and gender-based violence [43] hold promise; such programs seek to directly discuss and reconfigure gender roles in the direction of more gender equitable relationships [44]. Additionally, many gender inequalities exist at a structural level through cultural norms, laws, and institutions. Future research should also seek to implement structural-level interventions so that women may live in environments that better facilitate their control over their own sexual and reproductive health. Though structural-level interventions can be challenging both to implement and evaluate, they can have significant impact [45].

This is the first systematic review of interventions to improve self-efficacy and empowerment around safer sex and reproductive health decision-making for women living with HIV. The limitations of the existing evidence indicate a need for further research to determine the impact of empowerment and self-efficacy interventions. Future studies should include measurement of a wider range of sexual and reproductive health and rights outcomes, including both proximal empowerment and more distal health outcome measures. Studies should ensure the meaningful participation of the community of women living with HIV in study design. Interventions should also be explicit about how their content addresses unequal gender power relations. Such studies would allow for clear conclusions on how these types of interventions may improve the SRH of women living with HIV.
Supporting information
S1 Appendix. Full search strategy for PubMed.
(DOCX)
S1 Checklist. PRISMA checklist.
(DOCX)

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