CONSORT-EHEALTH (V 1.6.1) - Submission/Publication Form

The CONSORT-EHEALTH checklist is intended for authors of randomized trials evaluating web-based and Internet-based applications/interventions, including mobile interventions, electronic games (incl multiplayer games), social media, certain telehealth applications, and other interactive and/or networked electronic applications. Some of the items (e.g. all subitems under item 5 - description of the intervention) may also be applicable for other study designs.

The goal of the CONSORT EHEALTH checklist and guideline is to be

- a) a guide for reporting for authors of RCTs,
- b) to form a basis for appraisal of an ehealth trial (in terms of validity)

CONSORT-EHEALTH items/subitems are MANDATORY reporting items for studies published in the Journal of Medical Internet Research and other journals / scientific societies endorsing the checklist.

Items numbered 1., 2., 3., 4a., 4b etc are original CONSORT or CONSORT-NPT (non-pharmacologic treatment) items.

Items with Roman numerals (i., ii, iii, iv etc.) are CONSORT-EHEALTH extensions/clarifications.

As the CONSORT-EHEALTH checklist is still considered in a formative stage, we would ask that you also RATE ON A SCALE OF 1-5 how important/useful you feel each item is FOR THE PURPOSE OF THE CHECKLIST and reporting guideline (optional).

Mandatory reporting items are marked with a red *.

In the textboxes, either copy & paste the relevant sections from your manuscript into this form - please include any quotes from your manuscript in QUOTATION MARKS, or answer directly by providing additional information not in the manuscript, or elaborating on why the item was not relevant for this study.

YOUR ANSWERS WILL BE PUBLISHED AS A SUPPLEMENTARY FILE TO YOUR PUBLICATION IN JMIR AND ARE CONSIDERED PART OF YOUR PUBLICATION (IF ACCEPTED). Please fill in these questions diligently. Information will not be copyedited, so please use proper spelling and grammar, use correct capitalization, and avoid abbreviations.

DO NOT FORGET TO SAVE AS PDF _AND_ CLICK THE SUBMIT BUTTON SO YOUR ANSWERS ARE IN OUR DATABASE !!!

Citation Suggestion (if you append the pdf as Appendix we suggest to cite this paper in the caption):

Eysenbach G, CONSORT-EHEALTH Group

CONSORT-EHEALTH: Improving and Standardizing Evaluation Reports of Web-based and Mobile Health Interventions

J Med Internet Res 2011;13(4):e126

URL: http://www.jmir.org/2011/4/e126/

* Required Your name * First Last Lisa Dulli Primary Affiliation (short), City, Country * University of Toronto, Toronto, Canada Family Health International (FHI 360) Your e-mail address * abc@gmail.com Idulli@fhi360.org Title of your manuscript * Provide the (draft) title of your manuscript. SMART Connections: Results from a randomized, controlled trial of an online, social mediabased support group for youth living with HIV in Nigeria Name of your App/Software/Intervention * If there is a short and a long/alternate name, write the short name first and add the long name in brackets.

doi: 10.2196/jmir.1923

PMID: 22209829

SMART Connections

Evaluated Version (if any)
e.g. "V1", "Release 2017-03-01", "Version 2.0.27913"
Not applicable
Language(s) * What language is the intervention/app in? If multiple languages are available, separate by comma (e.g. "English, French")
English
URL of your Intervention Website or App
e.g. a direct link to the mobile app on app in appstore (itunes, Google Play), or URL of the website. If the intervention is a DVD or hardware, you can also link to an Amazon page.
Your answer
URL of an image/screenshot (optional)
Your answer
Accessibility *
Can an enduser access the intervention presently?
access is free and open
access only for special usergroups, not open
access is open to everyone, but requires payment/subscription/in-app purchases
app/intervention no longer accessible
Other:

Primary Medical Indication/Disease/Condition *
e.g. "Stress", "Diabetes", or define the target group in brackets after the condition, e.g. "Autism (Parents of children with)", "Alzheimers (Informal Caregivers of)"
HIV
Primary Outcomes measured in trial *
comma-separated list of primary outcomes reported in the trial
Retention in HIV treatment (care)
Secondary/other outcomes
Are there any other outcomes the intervention is expected to affect?
HIV-related knowledge, social support, adherence to anti-retroviral therapy
Recommended "Dose" *
What do the instructions for users say on how often the app should be used?
Approximately Daily
Approximately Weekly
Approximately Monthly
Approximately Yearly
as needed"
Other:

Approx. Percentage of Users (starters) still using the app as recommended after 3 months *	
unknown / not evaluated	
0-10%	
11-20%	
21-30%	
31-40%	
<u>41-50%</u>	
51-60%	
61-70%	
71%-80%	
81-90%	
91-100%	
Other:	
Overall, was the app/intervention effective? *	
yes: all primary outcomes were significantly better in intervention group vs control	
partly: SOME primary outcomes were significantly better in intervention group vs control	
ono statistically significant difference between control and intervention	
outcomes potentially harmful: control was significantly better than intervention in one or more	
inconclusive: more research is needed	
Other:	

Article Preparation Status/Stage * At which stage in your article preparation are you currently (at the time you fill in this form)
not submitted yet - in early draft status
o not submitted yet - in late draft status, just before submission
submitted to a journal but not reviewed yet
submitted to a journal and after receiving initial reviewer comments
submitted to a journal and accepted, but not published yet
O published
Other:
Journal *
If you already know where you will submit this paper (or if it is already submitted), please provide the journal name (if it is not JMIR, provide the journal name under "other") not submitted yet / unclear where I will submit this Journal of Medical Internet Research (JMIR) JMIR mHealth and UHealth JMIR Serious Games JMIR Mental Health JMIR Public Health JMIR Formative Research Other JMIR sister journal Other:

Is this a full powered effectiveness trial or a pilot/feasibility trial? *
O Pilot/feasibility
Fully powered
Manuscript tracking number *
If this is a JMIR submission, please provide the manuscript tracking number under "other" (The ms tracking number can be found in the submission acknowledgement email, or when you login as author in JMIR. If the paper is already published in JMIR, then the ms tracking number is the four-digit number at the end of the DOI, to be found at the bottom of each published article in JMIR)
o no ms number (yet) / not (yet) submitted to / published in JMIR
Other:
TITLE AND ABSTRACT
TITLE AND ABSTRACT
TITLE AND ABSTRACT 1a) TITLE: Identification as a randomized trial in the title
 1a) TITLE: Identification as a randomized trial in the title 1a) Does your paper address CONSORT item 1a? * 1.e does the title contain the phrase "Randomized Controlled Trial"? (if not, explain the reason under
 1a) TITLE: Identification as a randomized trial in the title 1a) Does your paper address CONSORT item 1a? * I.e does the title contain the phrase "Randomized Controlled Trial"? (if not, explain the reason under "other")

1a-i) Identify the mode of de	livery ir	n the title	9					
Identify the mode of delivery. Prefera title. Avoid ambiguous terms like "on includes non-web-based Internet com offline products are used. Use "virtua only in the context of "online support terms for the class of products (such	line", "virtonponents al" only in groups". as "mobi	ual", "inter (e.g. emai the contex Compleme	active". Us), use "cor t of "virtua ent or subs	se "Interne nputer-ba al reality" (stitute pro	t-based" o sed" or "ele (3-D worlds duct name	nly if Intervention ectronic" only if s). Use "online" s with broader		
application runs on different platforms.								
	1	2	3	4	5			
	·	_		•	· ·			
subitem not at all important	0	0	0	0	0	essential		
Does your paper address sul	bitem 1a	a-i? *						
Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly experience.	m manuso iuscript), o	cript title (or elaborat	e on this i	tem by pro	oviding add	litional		
	om a ran	domized	controll	ed trial o	f an onlin	e social media		
"SMART Connections: Results frobased support group for youth li	ving with	n HIV in N	igeria"					
"SMART Connections: Results fro based support group for youth li	ving with	or impo	igeria" rtant co	-interve	entions ir	n title		
"SMART Connections: Results frobased support group for youth living la-ii) Non-web-based components	ving with	or impo	igeria" rtant co	-interve	entions ir	n title		
"SMART Connections: Results frobased support group for youth living la-ii) Non-web-based components	onents	or impo	igeria" rtant co	-interve	entions ir any (e.g., "	n title		
"SMART Connections: Results frobased support group for youth living and a support group for youth living group group for youth living group gro	onents or import	or impo tant co-int	igeria" rtant co	-interve	entions ir any (e.g., "	n title with telephone		
"SMART Connections: Results frobased support group for youth living the support of the support o	onents of or important of the original of the	or impo tant co-int 2 2-ii? cript title (rtant coerventions 3 include que on this i	-interve	entions ir any (e.g., " 5	essential rks "like this" to		
"SMART Connections: Results frobased support group for youth living and paste relevant sections from the section in the section in the section is section."	onents of or important of the original of the	or impo tant co-int 2 2-ii? cript title (rtant coerventions 3 include que on this i	-interve	entions ir any (e.g., " 5	essential		

1a-iii) Primary condition or target group in the title Mention primary condition or target group in the title, if any (e.g., "for children with Type I Diabetes") Example: A Web-based and Mobile Intervention with Telephone Support for Children with Type I Diabetes: Randomized Controlled Trial										
	1 2 3 4 5									
subitem not at all important	0	0	0	0	0	essential				
Does your paper address sur Copy and paste relevant sections fro indicate direct quotes from your man information not in the ms, or briefly Yes "youth living with HIV in Nig	om manusc nuscript), c explain wh	cript title (i or elaborat	e on this i	tem by pro	viding add	litional				
1b) ABSTRACT: Structured conclusions NPT extension: Description of experstatus.		•								
1b-i) Key features/functional comparator in the METHOD Mention key features/functionalities possible, also mention theories and systematic reviewers and indexers by what the main paper is reporting. If adding it)	S sectio /compone principles y including	n of the ints of the used for d g importan	ABSTRA intervention esigning to t synonym	ACT on and cor he site. Ke ns. (Note: (mparator in ep in mind Only report	the abstract. If the needs of in the abstract				
	1	2	3	4	5					
subitem not at all important	0	0	0	0	0	essential				

"structured support group interve media platform"	ention, SI	MART Co	nnection	s deliver	ed throug	h a social-
1b-ii) Level of human involve	ment in	the ME	THODS	section	of the A	BSTRACT
Clarify the level of human involvemer "therapist/nurse/care provider/physic if any). (Note: Only report in the abstr from the main body of text, consider	cian-assis act what	ted" (men the main p	tion numb	er and exp	ertise of p	roviders involved,
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper address sub Copy and paste relevant sections from this" to indicate direct quotes from you information not in the ms, or briefly e	m the mar our manus	nuscript ab script), or e	elaborate d	on this ite	n by provic	ling additional
Copy and paste relevant sections from this to indicate direct quotes from your information not in the ms, or briefly e	m the mar our manus	nuscript ab script), or e	elaborate d	on this ite	n by provic	ling additional
Copy and paste relevant sections from this" to indicate direct quotes from your information not in the ms, or briefly e	m the mar our manus xplain wh	nuscript abscript), or e	elaborate d	on this iter	n by provic levant for y	ling additional rour study
Copy and paste relevant sections from this" to indicate direct quotes from yo	m the mar our manus xplain wh	nuscript abscript), or e y the item	elaborate d is not app	on this iter licable/re t) vs. fa	n by provic levant for y	ling additional rour study
Copy and paste relevant sections from this" to indicate direct quotes from your information not in the ms, or briefly export answer 1b-iii) Open vs. closed, web-assessments in the METHOD Mention how participants were recruiclinic or a closed online user group (of trial, or there were face-to-face compoutcomes were self-assessed throug traditional offline trials, an open trial researchers and participants know we "blinded" or "unblinded" to indicated to usually refers to "open access" (i.e. p	based (S section ited (online closed use onents (a) the question (open-labe hich treation the level of articipant	self-ass on of the evs. offlir ergroup tri s part of tl nnaires (as el trial) is ment is be of blinding s can self-	elaborate of is not appoint is common a type of common atype of common instead of enrol). (Note that is not appoint in the common in the c	t) vs. fa ACT om an ope arify if this ation or fo in web-ba dinical tria istered. To "open", a ote: Only r	ce-to-facen access was a pur rassessment sed trials) I in which I o avoid conserving the port in the	website or from a ely web-based ent). Clearly say if . Note: In both the ufusion, use web-based trials e abstract what
Copy and paste relevant sections from this" to indicate direct quotes from your information not in the ms, or briefly e Your answer 1b-iii) Open vs. closed, web-	based (S section ited (online closed use onents (a) the question (open-labe hich treation the level of articipant	self-ass on of the evs. offlir ergroup tri s part of tl nnaires (as el trial) is ment is be of blinding s can self-	elaborate of is not appoint is common a type of common atype of common instead of enrol). (Note that is not appoint in the common in the c	t) vs. fa ACT om an ope arify if this ation or fo in web-ba dinical tria istered. To "open", a ote: Only r	ce-to-facen access was a pur rassessment sed trials) I in which I o avoid conserving the port in the	website or from a ely web-based ent). Clearly say if . Note: In both the ufusion, use web-based trials e abstract what

Does your paper address subitem 1b-i? *

Copy and paste relevant sections fro this" to indicate direct quotes from y information not in the ms, or briefly e	our manus	script), or e	elaborate c	n this iter	n by provid	ling additional
"We collected data at baseline a extraction. We also conducted in participants."		_				
1b-iv) RESULTS section in ab	stract m	nust con	tain use	data		
Report number of participants enroll attrition/adherence metrics, use over outcomes. (Note: Only report in the amissing from the main body of text, or the main body	r time, nur abstract w	nber of log hat the ma	jins etc.), i	n addition	to primary	//secondary
	1	2	3	4	5	
	\bigcirc	\bigcirc	\bigcirc	0	0	essential
subitem not at all important	O					
Does your paper address su						
Does your paper address su Copy and paste relevant sections fro this" to indicate direct quotes from y	m the mai	nuscript ab script), or e	elaborate c	n this iter	n by provid	ling additional
Does your paper address su Copy and paste relevant sections fro	m the mai	nuscript ab script), or e	elaborate c	n this iter	n by provid	ling additional
Does your paper address su Copy and paste relevant sections fro this" to indicate direct quotes from y information not in the ms, or briefly e	m the mai our manus explain wh	nuscript abscript), or e	elaborate d is not app	on this iter licable/rel	n by provid evant for y	ling additional
Does your paper address su Copy and paste relevant sections fro this" to indicate direct quotes from y information not in the ms, or briefly e	SSION in the manusexplain when the sexplain when	nuscript abscript), or e y the item ive trials: I the interve iscuss reas	elaborate of is not appoint for ne Discuss the notion was sons. (Not	egative e primary not used, e: Only rep	trials outcome - discuss wl	ding additional vour study if the trial is hether negative abstract what the
Does your paper address su Copy and paste relevant sections fro this" to indicate direct quotes from y information not in the ms, or briefly e Your answer 1b-v) CONCLUSIONS/DISCU Conclusions/Discussions in abstract negative (primary outcome not changesults are attributable to lack of upt	SSION in the manusexplain when the sexplain when	nuscript abscript), or e y the item ive trials: I the interve iscuss reas	elaborate of is not appoint for ne Discuss the notion was sons. (Not	egative e primary not used, e: Only rep	trials outcome - discuss wl	ding additional vour study if the trial is hether negative abstract what the

Does your paper address subitem 1b-v?

Copy and paste relevant sections from the manuscript abstract (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"Our findings of improved HIV knowledge and high acceptability are encouraging, despite a lack of measurable effect on retention. Greater than anticipated retention in both groups was likely a result of external efforts that began part-way through the study."

INTRODUCTION

2a) In INTRODUCTION: Scientific background and explanation of rationale

2a-i) Problem and the type of system/solution

Describe the problem and the type of system/solution that is object of the study: intended as stand-alone intervention vs. incorporated in broader health care program? Intended for a particular patient population? Goals of the intervention, e.g., being more cost-effective to other interventions, replace or complement other solutions? (Note: Details about the intervention are provided in "Methods" under 5)

1 2 3 4 5

subitem not at all important O O O essential

Does your paper address subitem 2a-i? *

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"Despite a lack of age-disaggregated data on antiretroviral therapy (ART) coverage for youth, available data show that YLHIV enrolled in HIV care experience higher loss to follow up and suboptimal treatment adherence compared to younger children or adults."

"Developing and testing interventions to improve YLHIV outcomes in SSA is urgently needed and digital health interventions may present innovative opportunities to improve adherence and retention among youth in LMIC."

2a-ii) Scientific background, rationale: What is known about the (type of) system

Scientific background, rationale: What is known about the (type of) system that is the object of the study (be sure to discuss the use of similar systems for other conditions/diagnoses, if appropiate), motivation for the study, i.e. what are the reasons for and what is the context for this specific study, from which stakeholder viewpoint is the study performed, potential impact of findings [2]. Briefly justify the choice of the comparator.

	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

Does your paper address subitem 2a-ii? *

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"few interventions to improve these outcomes target youth specifically. Interventions to improve HIV-related outcomes implemented in low- and middle -income countries (LMIC) largely target adults and aim to improve ART adherence; fewer interventions target retention in care.[12-14] Support groups to improve health outcomes among people living with HIV (PLHIV), including YLHIV, are supported by some evidence to improve retention in care.[15-22]

Expanding access to mobile phone technology has increased interest in using this technology to improve health outcomes. Digital health interventions such as mobile reminders and interactive voice or SMS response have shown some effectiveness in improving adherence or retention among PLHIV in LMIC.[23-31] Though research has not examined these interventions specifically among YLHIV in LMIC, preliminary evidence in high income countries suggests such interventions are feasibility and may impact ART adherence on youth.[32-35] Furthermore, two recent studies (in South Africa and the US) integrated social networking into interventions for YLHIV to improve social support and found them to be acceptable and feasible.[36, 37]"

2b) In INTRODUCTION: Specific objectives or hypotheses

Does your paper address CONSORT subitem 2b? *

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"we set out to test the effectiveness of the SMART Connections intervention on HIV treatment retention among youth ages 15 to 24 years. We also examined SMART Connections effects on secondary outcomes of ART adherence, HIV knowledge, and social support."

METHODS

3a) Description of trial design (such as parallel, factorial) including allocation ratio

Does your paper address CONSORT subitem 3a? *

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"We conducted a 2-arm, parallel, randomized-controlled trial. Participants were individually randomized in a 1:1 ratio to either the SMART Connections intervention (intervention group) or standard of care (control group)."

3b) Important changes to methods after trial commencement (such as eligibility criteria), with reasons

Does your paper address CONSORT subitem 3b? *
Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study
"We originally planned to recruit YLHIV ages 15 to 22 years on ART for six months or less and expected to achieve our sample size within three months. Due to substantially lower than expected numbers of YLHIV enrolling in HIV treatment, we amended the study protocol and expanded the eligibility criteria to include YLHIV up to 24 years-old and on ART for 12 months or less. We also extended enrollment from three to eight months and added 3 facilities (originally 11)."
3b-i) Bug fixes, Downtimes, Content Changes
Bug fixes, Downtimes, Content Changes: ehealth systems are often dynamic systems. A description of changes to methods therefore also includes important changes made on the intervention or comparator during the trial (e.g., major bug fixes or changes in the functionality or content) (5-iii) and other "unexpected events" that may have influenced study design such as staff changes, system

failures/downtimes, etc. [2]. 1 2 3 4 5 subitem not at all important O O O o essential

Does your paper address subitem 3b-i?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Your answer

4a) Eligibility criteria for participants

Does your paper address CONSORT subitem 4a? *

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"YLHIV 15to 24 years-old and on ART for 12 months or less." and "In addition to age and ART eligibility criteria, participants had to demonstrate basic literacy for online chats, which was assessed at enrollment by asking the participant to read three short messages from intervention materials. We excluded individuals who knew they would be unable to attend the initial intervention group meeting if randomized to the intervention group, who were enrolled in another research study related to HIV retention or ART adherence, or who were severely ill and unable to provide informed consent at the time of recruitment."

4a-i) Computer / Internet literacy

Computer / Internet literacy is often an implicit "de facto" eligibility criterion - this should be explicitly clarified.

1 2 3 4 5

subitem not at all important

0 0

0

) essential

Does your paper address subitem 4a-i?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"...participants had to demonstrate basic literacy for online chats, which was assessed at enrollment by asking the participant to read three short messages from intervention materials."

4a-ii) Open vs. closed, web-based vs. face-to-face assessments: Open vs. closed, web-based vs. face-to-face assessments: Mention how participants were recruited (online vs. offline), e.g., from an open access website or from a clinic, and clarify if this was a purely web-based trial, or there were face-to-face components (as part of the intervention or for assessment), i.e., to what degree got the study team to know the participant. In online-only trials, clarify if participants were quasi-anonymous and whether having multiple identities was possible or whether technical or logistical measures (e.g., cookies, email confirmation, phone calls) were used to detect/prevent these.

subitem not at all important O O O O essential

Does your paper address subitem 4a-ii? *

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"We conducted face-to-face structured interviews with participants and extracted data from medical records at enrollment (baseline) and again at the completion of the intervention period (endline), approximately 6-9 months after enrollment. We also conducted in-depth interviews (IDI) at endline with two subsets of intervention group participants" "Eligible participants were sequentially recruited from patients attending clinic visits at study facilities." "Structured questionnaires programmed into password-protected computer tablets were used to collect data from participants at both baseline and endline. We also extracted medical record data (MRE) from patient charts at both time points into a separate form programmed into the tablets. Data from tablets were uploaded daily to a secure computer server. Trained interviewers administered a semi-structured IDI guide. IDIs were audio-recorded and transcribed verbatim."

4a-iii) Information giving during recruitment

Information given during recruitment. Specify how participants were briefed for recruitment and in the informed consent procedures (e.g., publish the informed consent documentation as appendix, see also item X26), as this information may have an effect on user self-selection, user expectation and may also bias results.

	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

Does your paper address sul Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e	m the mar uscript), c	nuscript (ir or elaborat	e on this i	tem by pro	oviding add	itional
Your answer						
4b) Settings and locations v	vhere th	ne data	were co	ollected		
Does your paper address CC Copy and paste relevant sections froi indicate direct quotes from your man information not in the ms, or briefly e "Health providers referred potent participate in the health facility." structured interviews and medica	m the mar uscript), c xplain wh tially elig We colled	nuscript (in or elaborat y the item ible patie cted data	nclude quo e on this in is not app ents to sto at basel	tem by pro licable/re udy staff	oviding add levant for y who recr	itional rour study uited them to
4b-i) Report if outcomes were (self-trials) or otherwise.				•	•	
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper address sub Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e Not applicable	m the mar uscript), c	nuscript (ir or elaborat	e on this i	tem by pro	oviding add	itional

4b-ii) Report how institution Report how institutional affiliations affiliations with prestigious hospitals regards to an intervention. (Not a require	are display s or univer	ed to pote sities may	ntial partic	cipants [or unteer rate	es, use, an	
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper address su	bitem 4	b-ii?				
Copy and paste relevant sections fro indicate direct quotes from your mar information not in the ms, or briefly e	om the mar nuscript), c	nuscript (ir or elaborat	e on this it	tem by pro	viding add	itional
Your answer						
5) The interventions for eac including how and when the	•				to allow	v replication,
5-i) Mention names, credent owners	tial, affili	ations o	f the de	veloper	s, spons	ors, and
Mention names, credential, affiliation are owners or developer of the softwomentioned elsewhere in the manuscr	are, this n					
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

Does your paper address subitem 5-i?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"Intervention Description

SMART Connections was informed through workshops conducted with stakeholders and YLHIV in Akwa Ibom State, Nigeria to gather input into design and content.[12, 13] The intervention was designed to promote retention in HIV care by leveraging social support and improving HIV-related knowledge and treatment literacy.[46, 47] Content of the structured support groups was adapted from an existing support group guide, Positive Connections and delivered through secret Facebook ™ groups.[48] The intervention was delivered over approximately 22 weeks (2 weeks per session) to groups of about 15-25 youth, with nearly daily activities (Figure 1).

Two community-based organizations were engaged to recruit support group facilitators and assist study staff in training them to deliver the intervention. Selected facilitators had received prior training to facilitate in-person support groups and were living with HIV themselves. Facilitators underwent a 5-day training and received an implementation guide, smart phone and a monthly data allowance. Facilitators met monthly with the study staff to debrief on challenges they'd encountered and provide support to one another. Once a sufficient number of participants was recruited and randomized to form a support group, each group began with an in-person meeting, during which participants met one another and the facilitator and agreed upon ground rules for participation. "

5-ii) Describe the history/development process

Describe the history/development process of the application and previous formative evaluations (e.g., focus groups, usability testing), as these will have an impact on adoption/use rates and help with interpreting results.

1 2 3 4 5
subitem not at all important O O O O essential

Does your paper address subitem 5-ii?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Your answer

Revisions and updating. Clearly ment (and comparator, if applicable) evaluation grocess, or whe Describe dynamic components such the replicability of the intervention (for	ated, or de ther the d as news f	escribe wh levelopme eeds or ch	ether the i nt and/or anging co	interventic content w intent whic	n underwe as "frozen"	nt major changes during the trial.
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper address sub	oitem 5	-iii?				
Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e	uscript), c	or elaborat	e on this i	tem by pro	viding add	itional
Your answer						
5-iv) Quality assurance meth Provide information on quality assura provided [1], if applicable.		ods to ens	sure accur	acy and q	uality of inf	ormation
	1	2	3	4	5	
subitem not at all important	1	2	3	4	5	essential
	0	0	3	4	5	essential
subitem not at all important Does your paper address subscriptions from the proper address from your man information not in the ms, or briefly expenses.	oitem 5-	-iv?	nclude quo	otes in quo	otation mar	ks "like this" to litional

screenshots/screen-capture used	video, a	and/or p	roviding	g flowch	arts of t	he algorithms
Ensure replicability by publishing the and/or providing flowcharts of the algorimciple be able to replicate the students.	gorithms ι	used. Repl	icability (i.	.e., other r		•
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper address sub	oitem 5-	-v?				
Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e	uscript), c	r elaborat	e on this i	tem by pro	oviding add	itional
Your answer						
5-vi) Digital preservation						
Digital preservation: Provide the URL disappear over the course of the year webcitation.org, and/or publishing th pages behind login screens cannot be without login.	s; also ma e source c	ake sure th code or sc	ne interver reenshots,	ntion is ard /videos al	chived (Inte	ernet Archive, e article). As
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper address sub	oitem 5-	-vi?				
Copy and paste relevant sections from		r elaborat	e on this i	tem by pro	oviding add	itional
indicate direct quotes from your man information not in the ms, or briefly e	xplain wh	y the item	is not app		-	,
indicate direct quotes from your man	xplain wh	y the item	іѕ посарр			·

Access: Describe how participants a (or were paid) or not, whether they had participants obtained "access to the editors/reviewers/readers, consider reviewers/readers to explore the approximation of the second control of	ad to be a platform a to provide	member o and Interne a "backdo	f specific et" [1]. To e or" login a	group. If kensure account or	nown, des cess for demo mod	cribe how de for
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper address su	bitem 5	-vii? *				
Copy and paste relevant sections fro indicate direct quotes from your mar information not in the ms, or briefly e	nuscript), d	or elaborat	e on this it	tem by pro	oviding add	litional
Your answer						
Your answer 5-viii) Mode of delivery, feat and comparator, and the the Describe mode of delivery, features/the theoretical framework [6] used to techniques, persuasive features, etc. description of the content (including how] it is tailored to individual circum feedback" [6]. This also includes a demediated communication is a composite [6]. It also includes information on proamount of text on pages, presence of	functional o design th , see e.g., where it is nstances a escription onent – wh	I framevities/complem (instru [7, 8] for to s coming for and allows of community of community of community of community of strategies	vork conents of actional strendingly rom and w users to t unication d nmunication es [1], inclu	the interverted the interverted the interverted their elivery chorn was synding page	ention and behaviour cludes an i ped it) [1], progress a annels and nchronous	comparator, and change n-depth whether [and ind receive – if computeror asynchronous
5-viii) Mode of delivery, feat and comparator, and the the Describe mode of delivery, features/t the theoretical framework [6] used to techniques, persuasive features, etc. description of the content (including how] it is tailored to individual circum feedback" [6]. This also includes a demediated communication is a compo [6]. It also includes information on pr	functional o design th , see e.g., where it is nstances a escription onent – wh	I framevities/complem (instru [7, 8] for to s coming for and allows of community of community of community of community of strategies	vork conents of actional strendingly rom and w users to t unication d nmunication es [1], inclu	the interverted the interverted the interverted their elivery chorn was synding page	ention and behaviour cludes an i ped it) [1], progress a annels and nchronous	comparator, and change n-depth whether [and ind receive – if computeror asynchronous

Does your paper address subitem 5-viii? *

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"Intervention Description

SMART Connections was informed through workshops conducted with stakeholders and YLHIV in Akwa Ibom State, Nigeria to gather input into design and content.[12, 13] The intervention was designed to promote retention in HIV care by leveraging social support and improving HIV-related knowledge and treatment literacy.[46, 47] Content of the structured support groups was adapted from an existing support group guide, Positive Connections and delivered through secret Facebook ™ groups.[48] The intervention was delivered over approximately 22 weeks (2 weeks per session) to groups of about 15-25 youth, with nearly daily activities (Figure 1).

Two community-based organizations were engaged to recruit support group facilitators and assist study staff in training them to deliver the intervention. Selected facilitators had received prior training to facilitate in-person support groups and were living with HIV themselves. Facilitators underwent a 5-day training and received an implementation guide, smart phone and a monthly data allowance. Facilitators met monthly with the study staff to debrief on challenges they'd encountered and provide support to one another.

Once a sufficient number of participants was recruited and randomized to form a support group, each group began with an in-person meeting, during which participants met one another and the facilitator and agreed upon ground rules for participation.

Standard of care services

All study participants continued to receive standard services available to YLHIV in study facilities and communities. Standard health services included: routine clinical care for HIV treatment including viral load tests; active case management by community volunteers with intensive adherence support during the first 4 weeks of ART; and enhanced adherence counseling for patients with unsuppressed viral loads.

All study participants received a smart phone (valued at US\$65) and a monthly data bundle (US\$3.5 equivalent to 1 gigabyte of data) for the duration of the intervention."

5-ix) Describe use parameters

Describe use parameters (e.g., intended "doses" and optimal timing for use). Clarify what instructions or recommendations were given to the user, e.g., regarding timing, frequency, heaviness of use, if any, or was the intervention used ad libitum.

	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

indicate direct quotes from your mar information not in the ms, or briefly e	nuscript), d	or elaborat	e on this i	tem by pro	oviding add	
Your answer						
5-x) Clarify the level of huma	an involv	vement				
Clarify the level of human involveme in the e-intervention or as co-interver as well as "type of assistance offere medium by which the assistance is chuman involvement required for the application outside of a RCT setting	ntion (deta d, the timin delivered". trial, and th	ail number ng and fre It may be he level of	and exper quency of necessary human in	tise of pro the suppo to disting volvement	fessionals rt, how it is uish betwe required f	involved, if any, in initiated, and the een the level of
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Copy and paste relevant sections fro indicate direct quotes from your mar information not in the ms, or briefly e	om the mar nuscript), c	nuscript (ii or elaborat	e on this i	tem by pro	oviding add	litional
Copy and paste relevant sections fro indicate direct quotes from your mar information not in the ms, or briefly of Your answer 5-xi) Report any prompts/relevant any prompts/relevant any prompts/reminders used use the application, what triggered the level of prompts/reminders required	om the man nuscript), c explain wh minders : Clarify if hem, frequ for the tria	nuscript (in or elaborat y the item s used there were lency etc.	e on this it is not app e prompts it may be r level of pr	tem by pro licable/re (letters, en necessary ompts/rei	nails, phor to distingu	litional your study ne calls, SMS) to nish between the
Does your paper address su Copy and paste relevant sections fro indicate direct quotes from your mar information not in the ms, or briefly of Your answer 5-xi) Report any prompts/rel Report any prompts/reminders used use the application, what triggered th level of prompts/reminders required application outside of a RCT setting	om the man nuscript), c explain wh minders : Clarify if hem, frequ for the tria	nuscript (in or elaborat y the item s used there were lency etc.	e on this it is not app e prompts it may be r level of pr	tem by pro licable/re (letters, en necessary ompts/rei	nails, phor to distingu	litional your study ne calls, SMS) to nish between the

Does your paper address sul	bitem 5	-xi? *				
Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly expressions and the ms are the companies.	uscript), c	or elaborat	e on this i	tem by pro	viding add	itional
Not applicable						
5-xii) Describe any co-interv			Ū			
Describe any co-interventions (incl. to addition to the targeted eHealth inter intervention. This includes training so the level of training required for the to RCT setting (discuss under item 21 –	vention, a essions an rial, and th	s ehealth ind support ne level of	interventio [1]. It may	on may not / be neces	be designessary to dis	ed as stand-alone tinguish between
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper address sul Copy and paste relevant sections froi indicate direct quotes from your man information not in the ms, or briefly e	m the mar luscript), c explain wh	nuscript (in or elaborat y the item	e on this it	tem by pro licable/rel	viding add evant for y	itional our study
were living with HIV themselves. implementation guide, smart phowith the study staff to debrief on another.	Facilitat	ors unde a monthly	rwent a 5 y data all	5-day traii owance.	ning and i Facilitato	received an rs met monthly
6a) Completely defined pre- measures, including how an	•	•	•		dary out	come

Does your paper address CONSORT subitem 6a? *

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"Retention: For our primary time-to-event analysis, we computed the time retained in care from study enrollment to the date the participant was no longer classified as active on treatment, consistent with PEPFAR indicator definitions.[50] We recorded dates for all scheduled clinic visits for each participant from study enrollment until the end of the study. If a participant failed to return after a scheduled visit for >28 days, the date of the missed visit was the date of loss to care recorded, unless a death or transfer of service was documented prior to the missed visit. For a small number of patients, their first missed scheduled visit was scheduled on or within 28 days before study enrollment. If the participant missed this first scheduled visit by >28 days, he/she was assigned a retention time of 0 (zero).

We also reported on treatment status at endline. To be considered active on treatment at endline, an individual must have attended a visit or had a follow-up visit scheduled within 28 days of the date of their endline questionnaire. For those who did not complete and endline questionnaire, an approximate endline date was used (based on the median time in the study of those enrolled the same month). Participants who died or transferred service were categorized accordingly."

Secondary outcome measures are contained in a table.

6a-i) Online questionnaires: describe if they were validated for online use and apply CHERRIES items to describe how the questionnaires were designed/deployed

If outcomes were obtained through online questionnaires, describe if they were validated for online use and apply CHERRIES items to describe how the questionnaires were designed/deployed [9].

	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

Does your paper address subitem 6a-i?

Copy and paste relevant sections from manuscript text

Not applicable

defined/measured/monitored	d					
Describe whether and how "use" (incl (logins, logfile analysis, etc.). Use/ad reported in any ehealth trial.						
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper address sub Copy and paste relevant sections from						
Your answer						
6a-iii) Describe whether, how was obtained Describe whether, how, and when qua emails, feedback forms, interviews, for	alitative fe	edback fro				· · · · · ·
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper address sub Copy and paste relevant sections from						

7b) When applicable, explan	ation o	f any in	terim aı	nalyses	and stop	oping
"The study was powered to detect retention at endline (0.45 in the corresponding to a hazard ratio of sided comparison using the log-reper study arm). Calculations also study follow-up."	ontrol g of 0.69, v ank test	roup and vith 80% . This res	0.575 in power ar sulted in	the inter nd 5% sig a total sa	vention g nificance mple size	roup), level for a two- e of 500 (250
Does your paper address sub Copy and paste relevant sections from indicate direct quotes from your mand information not in the ms, or briefly ex	n manusc uscript), c	cript title (or elaborat	e on this i	tem by pro	viding add	litional
subitem not at all important	0	0	0	0	0	essential
7a-i) Describe whether and h calculating the sample size Describe whether and how expected a			nto accou			
7a) How sample size was de NPT: When applicable, details of when addressed			ustering b	y care prov	rides or ce	nters was
Not applicable, no change to trial	outcom	nes				
Copy and paste relevant sections from indicate direct quotes from your manu information not in the ms, or briefly expenses.	uscript), c	or elaborat	e on this i	tem by pro	viding add	litional

Does your paper address CONSORT subitem 7b? *

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Not applicable

8a) Method used to generate the random allocation sequence

NPT: When applicable, how care providers were allocated to each trial group

Does your paper address CONSORT subitem 8a? *

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"The allocation sequence was generated using permuted blocks and stratified by local government areas (LGA), which are administrative sub-units of the state, by a biostatistician otherwise uninvolved in the study using a validated SAS macro - RANDOM (SAS version 9.4)."

8b) Type of randomisation; details of any restriction (such as blocking and block size)

Does your paper address CONSORT subitem 8b? *

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"The allocation sequence was generated using permuted blocks and stratified by local government areas (LGA), which are administrative sub-units of the state, by a biostatistician otherwise uninvolved in the study using a validated SAS macro - RANDOM (SAS version 9.4)."

9) Mechanism used to implement the random allocation sequence (such as sequentially numbered containers), describing any steps taken to conceal the sequence until interventions were assigned

indicate direct quotes from your man information not in the ms, or briefly e					_	
"Data collectors assigned partici Randomization groups were con Recruitment occurred at health f location for all facilities within a study staff or participants."	cealed ir acilities,	sequent but rand	tially num omizatio	nbered, s n was ma	ealed opa anaged fr	ique envelopes. om a central
10) Who generated the rand participants, and who assig			•			d
Does your paper address CC	ONSORT	subiter	m 10? *			
Copy and paste relevant sections fro indicate direct quotes from your man information not in the ms, or briefly e	uscript), d	or elaborat	e on this i	tem by pro	oviding add	litional
Yes, please see answers to previ	ious thre	e questic	ons.			
11a) If done, who was blinde participants, care providers NPT: Whether or not administering co	, those	assessi	ng outc	comes) a	and how	
11a-i) Specify who was blinde	ed, and	who wa	sn't			
Specify who was blinded, and who was participants [1, 3] (this should be cleassessors, those doing data analysis	arly ackno	wledged),	but it may	/ be possib	ole to blind	
		2	3	4	5	
	1					

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to

Does your paper address CONSORT subitem 9? *

		la-i? *				
Copy and paste relevant sections fro indicate direct quotes from your mar information not in the ms, or briefly e	nuscript), d	or elaborat	e on this i	tem by pro	oviding add	itional
No blinding of any subjects or study staff						
						<u>.</u>
11a-ii) Discuss e.g., whether 'intervention of interest" and						as the
nformed consent procedures (4a-ii) participants knew which intervention comparator".	can create	e biases aı	nd certain	expectation	ons - discu	
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Copy and paste relevant sections fro	m the mai	nuscript (i	-			
Copy and paste relevant sections fro ndicate direct quotes from your mar nformation not in the ms, or briefly e	m the mai	nuscript (i or elaborat	e on this i	tem by pro	oviding add	itional
Does your paper address surceptions from the properties of the pro	of the sehealth tria	nuscript (in or elaborately the item of the item similaritely als as it re	e on this it is not app	tem by pro licable/re	oviding add levant for y	litional rour study
Copy and paste relevant sections fro ndicate direct quotes from your mar nformation not in the ms, or briefly expour answer 11b) If relevant, description (this item is usually not relevant for exponential intervention to a active medication/intervention to a active medication/intervention paste relevant sections fro	of the sehealth triantervention	nuscript (in prelaborate of the item of t	e on this it is not app y of inter fers to sim n 11b? * nclude que	erventice in que	oviding add levant for y ons a placebo d	or sham
Copy and paste relevant sections fron dicate direct quotes from your marnformation not in the ms, or briefly expour answer [11b] If relevant, description (this item is usually not relevant for exponential intervention to a active medication/intervention to a active medication to a active medication to a active medication to active medication	of the sehealth triantervention	nuscript (in prelaborate of the item of th	y of inte fers to sim	erventice in quotes in quo	oviding add levant for y ons a placebo o	rks "like this" to

12a) Statistical methods used to compare groups for primary and secondary outcomes

NPT: When applicable, details of whether and how the clustering by care providers or centers was addressed

Does your paper address CONSORT subitem 12a? *

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"All analyses used an intent-to-treat (ITT) approach. The primary hypothesis was: YLHIV enrolled in HIV treatment services who participate in SMART Connections will be more likely to be retained in HIV care than YLHIV enrolled in HIV treatment services who do not participate in the intervention.

Kaplan-Meier cumulative retention probabilities are reported with 95% confidence intervals and plotted by study arm. Participants who were confirmed to have died or transferred to a facility outside the study facilities and for whom retention data could not be obtained were considered censored. The retention probabilities between the groups were compared with a log-rank test stratified by LGA with a two-sided alpha = 0.05. We also report on retention descriptively, examining lapses in care and return to treatment over the course of study follow-up.

The relationships between treatment exposure and secondary outcomes (ART adherence, HIV knowledge, and social support) as well as social isolation, depression, and HIV-related stigma were explored with t-tests for continuous outcomes and chi-square test for categorical outcomes, using two-sided tests, with significance level of 0.05."

12a-i) Imputation techniques to deal with attrition / missing values

Imputation techniques to deal with attrition / missing values: Not all participants will use the intervention/comparator as intended and attrition is typically high in ehealth trials. Specify how participants who did not use the application or dropped out from the trial were treated in the statistical analysis (a complete case analysis is strongly discouraged, and simple imputation techniques such as LOCF may also be problematic [4]).

	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

	0110111 12	2a-i? *					
Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e	uscript), c	or elaborat	e on this i	tem by pro	oviding add	itional	
Missing values are reported. No imputation was used to replace missing data.							
12b) Methods for additional analyses	analyse	es, such	as sub	group a	nalyses	and adjusted	
Does your paper address CC	ONSORT	「subiter	n 12b? *				
Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly expressions of the sections of the section of the	uscript), c	or elaborat	e on this i	tem by pro	oviding add	itional	
Not applicable.							
					_		
X26) REB/IRB Approval and I subheading under "Methods X26-i) Comment on ethics co	s"] (not	a CONS	SORT ite		ımended	d as	
subheading under "Methods	s"] (not	a CONS	SORT ite	em)	imended	d as	
subheading under "Methods	s"] (not ommitte	a CONS	SORT ite	em)		d as essential	
subheading under "Methods X26-i) Comment on ethics co	ommitte	ee appro	SORT ite	em)			
x26-i) Comment on ethics co	ommitte 1 Oitem X m the mar uscript), o	ee appro 2 2 26-i? nuscript (in	oval 3 onclude quoe on this i	4 Ottes in quotem by pro	otation man	essential ks "like this" to itional	

consent documents.	nucu (see	, 12,. 30.	e [6] for so	me items		
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper address suk	oitem X	26-ii?				
Copy and paste relevant sections fror indicate direct quotes from your man information not in the ms, or briefly e	uscript), c	or elaborat	e on this i	tem by pro	viding add	itional
Your answer						
						and the liberty
Safety and security procedures, incl.	privacy co and traini	onsideratio ng, availab	oility of a h	otline)		uce the likelihood
Safety and security procedures, incl.	privacy co	onsideratio		•	ken to red 5	uce the likelihood
X26-iii) Safety and security p Safety and security procedures, incl. p or detection of harm (e.g., education subitem not at all important	privacy co and traini	onsideratio ng, availab	oility of a h	otline)		uce the likelihood essential
Safety and security procedures, incl. por detection of harm (e.g., education	privacy co and traini	onsiderationg, availab	oility of a h	otline)		
Safety and security procedures, incl. por detection of harm (e.g., education subitem not at all important	privacy co and traini 1 O oitem X m the mar uscript), o	onsiderationg, available 2 26-iii? nuscript (in prelaborat	anclude quo	otline) 4 Outline otes in quotem by pro	otation man	essential eks "like this" to itional
Safety and security procedures, incl. por detection of harm (e.g., education subitem not at all important Does your paper address subscriptions and paste relevant sections from indicate direct quotes from your manifest.	privacy co and traini 1 O oitem X m the mar uscript), o	onsiderationg, available 2 26-iii? nuscript (in prelaborat	anclude quo	otline) 4 Outline otes in quotem by pro	otation man	essential eks "like this" to itional

13a) For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analysed for the primary outcome

NPT: The number of care providers or centers performing the intervention in each group and the number of patients treated by each care provider in each center

Does your paper address CONSORT subitem 13a? *

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Yes through a Consort flow diagram

13b) For each group, losses and exclusions after randomisation, together with reasons

Does your paper address CONSORT subitem 13b? (NOTE: Preferably, this is shown in a CONSORT flow diagram) *

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

We used an intent to treat analysis. Four participants were excluded as ineligible - they provided false eligibility information at enrollment.

13b-i) Attrition diagram

Strongly recommended: An attrition diagram (e.g., proportion of participants still logging in or using the intervention/comparator in each group plotted over time, similar to a survival curve) or other figures or tables demonstrating usage/dose/engagement.

	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essentia

Does your paper address subitem 13b-i? Copy and paste relevant sections from the manuscript or cite the figure number if applicable (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study Your answer 14a) Dates defining the periods of recruitment and follow-up

Does your paper address CONSORT subitem 14a? *

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Yes, dates are included. "We recruited 356 youth between September 2018 and April 2019, 353 (99.2%) of whom enrolled in the study (Figure 2). Four participants were removed after enrollment, having provided false eligibility information and deemed ineligible. Participants were then randomly allocated to the intervention group (n=177) or control group (n=172). At endline (June to November 2019), 108 participants were lost to follow-up from the study, including 4 who died, 9 who discontinued study participation and 95 who were not reachable for an endline interview. Endline interviews and MRE data were completed for 241 participants. MRE data were collected for an additional 84 participants who were not reachable for an endline interview but who had not elected to end study participation. We conducted IDIs with 21 intervention participants following the endline survey. "

14a-i) Indicate if critical "secular events" fell into the study period Indicate if critical "secular events" fell into the study period, e.g., significant changes in Internet resources available or "changes in computer hardware or Internet delivery resources" 1 2 3 4 5 subitem not at all important O O O O essential

Does your paper address subitem 14a-i?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"The context of HIV service delivery also changed during study implementation. A dedicated "surge" in PEPFAR-supported HIV services aimed to increase the number of PLHIV enrolled on treatment and retained in care began in study sites part way through the study.[69] This surge entailed intensive efforts to better support HIV treatment services and PLHIV in their care and treatment. Strategies including community-based ART initiation, multi-month ART dispensing for stable patients, and community-delivered ART refills were implemented to increase ART initiation and to improve treatment adherence and retention.[69] These changes meant that the average time between clinic visits and the number of visits participants had during the study varied. Although participants in both study arms were exposed to this changing context, retention increased dramatically during the surge; retention at endline was considerably higher in both study arms compared to retention rates indicated by the pre-surge programmatic data used to calculate effect size estimates. These external efforts may have masked any possible impact that the intervention had on retention."

14b) Why the trial ended or was stopped (early)

Does your paper address CONSORT subitem 14b? *

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Lastly, we intended to collect outcome data from participants after the intervention was completed, and at 1 year from enrollment (approximately 6 months later), but were unable to collect a second round of outcome data because of prolonged recruitment – the funding project came to an end before the final round of data collection could take place.

15) A table showing baseline demographic and clinical characteristics for each group

NPT: When applicable, a description of care providers (case volume, qualification, expertise, etc.) and centers (volume) in each group

Does your paper address CC		Subitei	11 10.			
Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e	uscript), c	or elaborat	e on this i	tem by pro	viding add	litional
Yes, Table 2.						
15-i) Report demographics a	ssociate	ed with	digital d	ivide iss	sues	
In ehealth trials it is particularly impo such as age, education, gender, socia participants, if known.		-			_	
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper address sul	oitem 15	5-i? *				
Copy and paste relevant sections from indicate direct quotes from your man	m the mar uscript), c	nuscript (ir or elaborat	e on this i	tem by pro	viding add	litional
Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e	m the mar uscript), c	nuscript (ir or elaborat	e on this i	tem by pro	viding add	litional
Copy and paste relevant sections froi indicate direct quotes from your man information not in the ms, or briefly e	m the mar uscript), c xplain wh	nuscript (ir or elaborat y the item	e on this i	tem by pro licable/re	oviding add	litional vour study
Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e lncluded in table 2. 16) For each group, number	m the mar uscript), c explain wh	nuscript (ir or elaborat y the item	e on this it is not app	tem by pro licable/re	oviding add levant for y	litional your study ed in each
Does your paper address sub Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e Included in table 2. 16) For each group, number analysis and whether the an	of part	nuscript (in or elaborat y the item	e on this it is not app s (denor original a	minator	oviding add levant for y include d groups	litional your study ed in each
Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly elementary and paste relevant sections from your man information not in the ms, or briefly elementary and pasted in table 2. 16) For each group, number analysis and whether the analy	of part nalysis v inators' provide de lds" [1], e.nts "used"	ruscript (in or elaboraty the item cicipants vas by continuitions: Fig., N exposithe interv	e on this it is not apprise (denor priginal acceptance).	minator assigne efinition (and effects asented, N	oviding add levant for y include d groups s et sizes) "a used more at specific	ed in each s cross a range of a than x times, N pre-defined time
Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly elementary and the control of the control o	of part nalysis v inators' provide de lds" [1], e.nts "used"	ruscript (in or elaboraty the item cicipants vas by continuitions: Fig., N exposithe interv	e on this it is not app is (denor original a ovide de Report N's sed, N con ention/con proup). Alw	minator efinition (and effect assigned)	oviding add levant for y include d groups s et sizes) "a used more at specific	ed in each s cross a range of a than x times, N pre-defined time

Does your paper address sub	oitem 16	5-i? *				
Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e	uscript), d	or elaborat	e on this i	tem by pro	oviding add	litional
Yes.						
16-ii) Primary analysis should	l be inte	ent-to-tr	eat			
Primary analysis should be intent-to-t the appropriate caveats that this is no		-	-			only "users", with
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper address subscriptions from the properties of the control of the c	m the mar uscript), c	nuscript (ii or elaborat	e on this i	tem by pro	oviding add	litional
17a) For each primary and se estimated effect size and its		•			•	•
Does your paper address CC	NIS O DI	Subiter	n 17a2 *			
Copy and paste relevant sections froi indicate direct quotes from your man information not in the ms, or briefly e	m the mar uscript), c	nuscript (ii or elaborat	nclude quo e on this i	otes in quo tem by pro	oviding add	litional
Yes.						

0	0	0	\bigcirc	\bigcirc	
				O	essential
the man script), o	uscript (ir r elaborat	e on this it	em by pro	viding add	itional
esenta	ition of	both ab	SOIUTE	and rela	пуе еттест
ISORT	subiter	n 17b? *			
script), o	r elaborat	e on this it	em by pro	viding add	itional
(see tal	ble 5)				
	sthe man script), o lain why esenta SSORT the man script), o lain why	escript), or elaborate plain why the item escript of the secretary of the manuscript (in script), or elaborate places are secretary of the manuscript (in script), or elaborate places are secretary of the manuscript (in script), or elaborate places are secretary or elaborate places are secretary or elaborate places.	the manuscript (include quoscript), or elaborate on this it plain why the item is not appose esentation of both above the manuscript (include quoscript), or elaborate on this it plain why the item is not appose it in the manuscript (include quoscript), or elaborate on this it plain why the item is not appose it in the manuscript (include quoscript).	the manuscript (include quotes in quoteript), or elaborate on this item by problain why the item is not applicable/relesentation of both absolute a series as a series of the manuscript (include quotes in quoteript), or elaborate on this item by problain why the item is not applicable/relesentation.	the manuscript (include quotes in quotation mar script), or elaborate on this item by providing add plain why the item is not applicable/relevant for your seemants of both absolute and related as a script). SORT subitem 17b? * the manuscript (include quotes in quotation mar script), or elaborate on this item by providing add plain why the item is not applicable/relevant for you seemants.

17a-i) Presentation of process outcomes such as metrics of use and intensity of

Yes						
18-i) Subgroup analysis of co	mparin	a only u	sers			
A subgroup analysis of comparing on stressed that this is a self-selected sa (see 16-iii).	ly users i	s not unco	mmon in e			
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Copy and paste relevant sections from	m the mar uscript), c	nuscript (ii or elaborat	e on this i	tem by pro	oviding add	itional
Does your paper address sub Copy and paste relevant sections fror Indicate direct quotes from your mani Information not in the ms, or briefly ex Your answer	m the mar uscript), c	nuscript (ii or elaborat	e on this i	tem by pro	oviding add	itional
Copy and paste relevant sections from ndicate direct quotes from your mand nformation not in the ms, or briefly ex	n the mai uscript), c xplain wh	nuscript (in or elaborat y the item	e on this i	tem by pro licable/re	oviding add levant for y	itional
Copy and paste relevant sections from ndicate direct quotes from your maninformation not in the ms, or briefly export answer (Our answer (19) All important harms or unifor specific guidance see CONSORT	n the mar uscript), c xplain wh nintenc for harms	nuscript (in or elaborat y the item ded effe	e on this it	tem by pro licable/re	oviding add levant for y	itional
Copy and paste relevant sections from ndicate direct quotes from your maninformation not in the ms, or briefly exfour answer Your answer 19) All important harms or unity	n the manuscript), oxplain when the manuscript) on SOR1 m the manuscript), oxplain the manuscript the	nuscript (in prelaborate of the laborate of th	e on this it is not appoint to the content of the c	ach gro	up otation man	itional rour study ks "like this" to itional

19-i) Include privacy breaches, technical problems Include privacy breaches, technical problems. This does not only include physical "harm" to participants, but also incidents such as perceived or real privacy breaches [1], technical problems, and other unexpected/unintended incidents. "Unintended effects" also includes unintended positive effects [2].								
	1	2	3	4	5			
subitem not at all important	0	0	0	0	0	essential		
Does your paper address su Copy and paste relevant sections fro indicate direct quotes from your mar information not in the ms, or briefly e	m the mai	nuscript (in or elaborat	e on this it	em by pro	viding add	litional		
19-ii) Include qualitative feed staff/researchers Include qualitative feedback from pa strengths and shortcomings of the a or uses. This includes (if available) reby the developers.	rticipants pplication	or observa , especiall	ations fron	n staff/res oint to unii	searchers, i	if available, on nexpected effects		
	1	2	3	4	5			
subitem not at all important	0	0	0	0	0	essential		

	your paper			o+ (in ald	too in	no orbe IIII 1	sio" +-
indicate	e direct quotes	nt sections from from your manus ms, or briefly exp	script), or elab	orate on this it	em by providing	g additional	

participants, including those with low intervention participation, stated they received social support from facilitators and other group members. These participants described receiving encouragement and advice, having people to "share my feelings" with or "someone to talk to," and receiving answers to factual and personal questions. One participant described: I felt like I was not alone in the journey and it was really cool ... it was amazing. I don't know how to say it in words but it's something to build (us up) because sometimes we can't just do it by ourselves. We need to find people in the situations with us for us get stronger, so the group actually made me stronger. -19-year-old female, low participation Some support was related to self-management, such as encouragement to adhere to ART. For example, one participant described group members as "...people that would encourage you no matter anything, they tell you no matter anything, that they're okay with the drugs, ... (they) encourage you to take the ART" (22-year-old male, low participation). Participants frequently received multiple types of support and often supported other group members by sharing their own experiences or providing emotional support and advice. Nearly all IDI participants stated they felt a sense of connectedness with the group, sometimes described like "a family" or that participants got along "as brother and sister." This feeling was often attributed to having group members of the same age range and HIV status and conferred a sense of safety and confidentiality within the group. In endline questionnaires, when asked what they liked most about the intervention, the most common responses included receiving encouragement and support, ability to share their problems, and feeling a sense of unity or belonging.

Yes. "Although the two study arms did not differ significantly on social support, nearly all IDI

When asked about sources of social support outside of the intervention group, most said they didn't receive social support outside of the group, sometimes elaborating that they preferred not to disclose their HIV status to family and friends due to fear of stigma:

I don't like disclosing... I don't know if that person is a victim of HIV...the person may start broadcasting me. Things like that so that's why I don't tell her, I didn't tell people... They will not be able to encourage me since they don't know what I'm passing through ... I keep it (my status) to myself.

-19-year-old male, high participation

Only a couple of IDI participants stated they received support from family members; a few mentioned a healthcare provider."

"IDI participants often conveyed appreciation for learning about practical aspects of managing HIV, such as taking ART at consistent times during the day and eating "a balanced diet" to support overall health. Many participants also liked learning why medication adherence is important. Participants recalled learning that ART adherence would help them to feel healthier and achieve a longer life. One participant described, "by taking the drugs and eating your food every day, your body will be okay ... but if you avoid taking that drugs and you are not doing anything, you don't go for test, you might die at any point in time and nobody will know the purpose of your death" (22-year-old female, high participation). Many IDI participants also expressed enthusiasm for the social and interactive elements of the intervention, such as riddles posed by the facilitator. Most felt the Facebook platform was acceptable, reporting that the "secret" groups ensured their privacy and the online format allowed them to interact with the group at their convenience. One participant described, "you'll use it [the group] at the comfort of your home, not pressing you to go out, do this, every time to hear about the new information that you need to learn. Peacefully, you'll just learn inside your room, inside your, the comfort of your own home. So, I was very happy about that one" (17-year-old male, high participation). "

22) Interpretation consistent considering other relevant of NPT: In addition, take into account the expertise of care providers or centers.	evidenc e choice (e of the com				
22-i) Restate study questions starting with primary outcon Restate study questions and summar outcomes and process outcomes (us	nes and	process	s outcor	nes (use	e)	·
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Copy and paste relevant sections from			-			
indicate direct quotes from your man information not in the ms, or briefly e	. ,				-	
·	ention wa knowled improve ntly. IDIs Interven	y the item as design ge and so retentior data pro ition parti	ed to imp cial supp or socia vided evic	prove treated or the control of the	evant for y atment re findings i t; howeve perceive whelming	tention among ndicate the r, HIV-related d
"The SMART Connections interver YLHIV by improving HIV-related lintervention did not significantly knowledge did improve signification improvements in social support.	ention waknowledge improvently. IDIs Intervente online p	y the item as design ge and so retention data pro ation parti clatform,	ed to importial supportions and repo	prove treatort. Our lasuppor dence of also over rted it he	evant for y atment re findings i t; howeve perceive whelming id helped	tention among ndicate the r, HIV-related d ily found the them in their
"The SMART Connections intervery YLHIV by improving HIV-related lintervention did not significantly knowledge did improve signification improvements in social support. intervention acceptable, liked the HIV treatment."	ention waknowledge improvently. IDIs Intervente online p	y the item as design ge and so retention data pro ation parti clatform,	ed to importial supportions and repo	prove treatort. Our lasuppor dence of also over rted it he	evant for y atment re findings i t; howeve perceive whelming id helped	tention among ndicate the r, HIV-related d ily found the them in their

Does your paper address subitem 22-ii?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"Most study participants were over 19 years-old and, although much of the content is relevant to the people of all ages, we believe future research should exploring whether some of the intervention content should be tailored to the differing developmental needs of those 15-19 years and those 20-24 years."

"Future research should include viral load as a primary outcome for interventions designed to improve health outcomes among PLHIV."

"One possible reason for the lack of an intervention effect on social support may have been that the measure we used was not HIV-specific; we may not have captured the type of social support participants felt they received through the intervention. Given challenges with regard to stigma and disclosure, further research is needed to operationalize and measure social support associated with HIV."

"Moving forward, we suggest a few adaptations to the SMART Connections intervention and to continue to examine its potential effects. Expanding groups to include YLHIV on ART for >1 year may provide better support to those newly initiating treatment, as well help meet their own informational and support needs. Findings from IDIs suggest the intervention had a perceived effect on social support. We believe the tool we used to measure social support may not have adequately distinguished between more general social support and social support related to HIV. Further work to develop and test measures of social support that better reflect the support given/provided in the context of HIV should be pursued. Lastly, focusing on more reliable outcome measures, such as viral load, is strongly recommended for interventions attempting to improve HIV-related health outcomes."

20) Trial limitations, addressing sources of potential bias, imprecision, and, if relevant, multiplicity of analyses

20-i) Typical limitations in ehealth trials

Typical limitations in ehealth trials: Participants in ehealth trials are rarely blinded. Ehealth trials often look at a multiplicity of outcomes, increasing risk for a Type I error. Discuss biases due to non-use of the intervention/usability issues, biases through informed consent procedures, unexpected events.

	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

Does your paper address subitem 20-i? *

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"Limitations

Despite many strengths, including the use of a rigorous, experimental study design, the study had a number of limitations. Several factors limited study implementation and perhaps the interpretation of results. Results may not be generalizable to other youth outside our study areas. Secondly, recruitment lagged, taking more than twice as long as planned because YLHIV enrollment in ART was substantially lower than estimated using HIV service data before the study. Despite protocol modifications, we were only able to enroll 349 (70%) of the planned 500 participants, reducing the study's power. Slow enrollment also prolonged the time necessary to enroll enough intervention participants to form a support group in many cases. Thus, participants contributed different amounts of time to the study and some intervention group participants waited months before their support group could begin. This perhaps added to their risk of loss to follow up before initiating the intervention."

21) Generalisability (external validity, applicability) of the trial findings

NPT: External validity of the trial findings according to the intervention, comparators, patients, and care providers or centers involved in the trial

21-i) Generalizability to other populations

Generalizability to other populations: In particular, discuss generalizability to a general Internet population, outside of a RCT setting, and general patient population, including applicability of the study results for other organizations

1 2 3 4 5
subitem not at all important O O O O essential

Does your paper address subitem 21-i?

Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Your answer

routine application setting Discuss if there were elements in the prompts/reminders, more human inv impact the omission of these elements applied outside of a RCT setting.	olvement,	training se	essions or	other co-i	nterventior	ns) and what
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper address sull Copy and paste relevant sections fro indicate direct quotes from your man information not in the ms, or briefly e	m the mar	nuscript (ir or elaborat	e on this it	tem by pro	viding add	itional
Your answer						
OTHER INFORMATION						
23) Registration number and	d name	of trial	registry	,		
Does your paper address CC Copy and paste relevant sections fro indicate direct quotes from your man information not in the ms, or briefly e	m the mar	nuscript (ir or elaborat	nclude quo e on this it	tem by pro	viding add	itional
ClinicalTrials.gov NCT03516318		,				

21-ii) Discuss if there were elements in the RCT that would be different in a

Does your paper address CC Cite a Multimedia Appendix, other ref (include quotes in quotation marks "li elaborate on this item by providing ac not applicable/relevant for your study The full protocol has not been pu	erence, o ke this" to Iditional i	r copy and o indicate nformation	paste rele direct quot	tes from y	our manus	cript), or
25) Sources of funding and of funders	other s	upport ((such as	supply	of drug	s), role of
Does your paper address CC	NSORT	「subiter	n 25? *			
Copy and paste relevant sections from indicate direct quotes from your man- information not in the ms, or briefly e	uscript), d	or elaborat	e on this it	em by pro	viding add	itional
"This study was funded by the US Relief, under task order contract under IDIQ contract number AID- contents of this publication are t reflect the views of the USAID or	number OAA-I-15 he sole r	AID-OAA 5-00009, ` responsib	-TO-15-00 YouthPov oility of Fl	0003 and ver: Implo HI 360 ar	YouthPo ementation	wer Action, on. The
X27) Conflicts of Interest (ne	ot a CC	ONSORT	item)			
X27-i) State the relation of th	e study	/ team to	owards t	the syst	em bein	g evaluated
In addition to the usual declaration of study team towards the system being identical with the developers/sponsor	evaluate	d, i.e., stat	e if the au	•		
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential

Does your ba	per address subitem X27-i?
indicate direct qu	elevant sections from the manuscript (include quotes in quotation marks "like this" to notes from your manuscript), or elaborate on this item by providing additional in the ms, or briefly explain why the item is not applicable/relevant for your study
Your answer	
About the Co	ONSORT EHEALTH checklist
As a result of	using this checklist, did you make changes in your manuscript? *
yes, major	changes
yes, minor	changes
O no	
What were the checklist?	ne most important changes you made as a result of using this
Small details o	f things already included.
	me did you spend on going through the checklist INCLUDING ges in your manuscript *

As a result of using this checklist, do you think your manuscript has improved? *
yes
O no
Other:
Would you like to become involved in the CONSORT EHEALTH group?
This would involve for example becoming involved in participating in a workshop and writing an "Explanation and Elaboration" document
O yes
o no
Other:
Any other comments or questions on CONSORT EHEALTH
The checklist has a number of redundant questions. Copying and pasting sections of the paper seems a lot of work for a review to review.
STOP - Save this form as PDF before you click submit To generate a record that you filled in this form, we recommend to generate a PDF of this page (on a Mac, simply select "print" and then select "print as PDF") before you submit it.
When you submit your (revised) paper to JMIR, please upload the PDF as supplementary file.
Don't worry if some text in the textboxes is cut off, as we still have the complete information in our database. Thank you!
Final step: Click submit! Click submit so we have your answers in our database!

Submit

Google Forms