# 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 (nonpharmacologic 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

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: <a href="http://www.jmir.org/2011/4/e126/">http://www.jmir.org/2011/4/e126/</a>

doi: 10.2196/jmir.1923 PMID: 22209829

mebetz@gmail.com (not shared) Switch account



Draft saved

\* Required

Your name \* First Last Marian Betz Primary Affiliation (short), City, Country \* University of Toronto, Toronto, Canada University of Colorado Anschutz Medical Camı Your e-mail address \* abc@gmail.com marian.betz@cuanschutz.edu Title of your manuscript \* Provide the (draft) title of your manuscript. The Safe at Home Study: Protocol for a Randomized Controlled Trial of a Web-Based Decision Aid for Caregivers of Persons with Dementia with Firearm Access 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. Safety in Dementia (SiD) Evaluated Version (if any) e.g. "V1", "Release 2017-03-01", "Version 2.0.27913" Your answer Language(s) \* What language is the intervention/app in? If multiple languages are available, separate by comma (e.g. "English, French") English, Spanish

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. https://safetyindementia.org/ 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)" Alzheimers (Informal Caregivers of) Primary Outcomes measured in trial \* comma-separated list of primary outcomes reported in the trial preparation for decision making Secondary/other outcomes Are there any other outcomes the intervention is expected to affect? firearm access, decision quality, caregiver burden

Approximately Daily Approximately Weekly Approximately Monthly Approximately Yearly  a "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% 21-30% 31-40% 41-50% 51-60% 61-70% 71%-80% 91-100% 0 ther:		ommended "Dose" * t do the instructions for users say on how often the app should be used?
Approximately Monthly Approximately Yearly  as needed" Other:  Approx. Percentage of Users (starters) still using the app as recommended after * 3 months  unknown / not evaluated O-10% 11-20% 21-30% 21-30% 31-40% 41-50% 51-60% 61-70% 71%-80% 91-100%	0	Approximately Daily
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<ul><li>81-90%</li><li>91-100%</li></ul>	0	61-70%
91-100%	0	71%-80%
	0	81-90%
Other:	0	91-100%
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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
on statistically significant difference between control and intervention
O potentially harmful: control was significantly better than intervention in one or more outcomes
inconclusive: more research is needed
Other: study protocol, work ongoing
Article Preparation Status/Stage *  At which stage in your article preparation are you currently (at the time you fill in this form)
onot submitted yet - in early draft status
onot 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")
onot 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

Is this a full powered effective Pilot/feasibility Fully powered	eness tr	ial or a p	oilot/feas	sibility t	rial? *				
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TITLE AND ABSTRACT									
1a) TITLE: Identification as a r	andom	iized tria	l in the t	itle					
1a) Does your paper address I.e does the title contain the phrareason under "other")  • yes  Other:				ed Trial"?	' (if not, e	xplain the			
Identify the mode of delivery. Pre "electronic game" in the title. Avo Use "Internet-based" only if Inter email), use "computer-based" or only in the context of "virtual rea support groups". Complement of	1a-i) Identify the mode of delivery in the title  Identify the mode of delivery. Preferably use "web-based" and/or "mobile" and/or "electronic game" in the title. Avoid ambiguous terms like "online", "virtual", "interactive".  Use "Internet-based" only if Intervention includes non-web-based Internet components (e.g. email), use "computer-based" or "electronic" only if offline products are used. Use "virtual" only in the context of "virtual reality" (3-D worlds). Use "online" only in the context of "online support groups". Complement or substitute product names with broader terms for the class of products (such as "mobile" or "smart phone" instead of "iphone"), especially if the application runs on different platforms								
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subitem not at all important	0	0	0	•	0	essential			
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Does your paper address subitem 1a-i? \* Copy and paste relevant sections from manuscript title (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 Safe at Home Study: Protocol for a Randomized Controlled Trial of a Web-Based Decision Aid for Caregivers of Persons with Dementia with Firearm Access" 1a-ii) Non-web-based components or important co-interventions in title Mention non-web-based components or important co-interventions in title, if any (e.g., "with telephone support"). 2 3 5 1 essential subitem not at all important Clear selection Does your paper address subitem 1a-ii? Copy and paste relevant sections from manuscript title (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 Safe at Home Study: Protocol for a Randomized Controlled Trial of a Web-Based Decision Aid for Caregivers of Persons with Dementia with Firearm Access" 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 5  $\circ$  $\circ$  $\circ$ • subitem not at all important essential Clear selection

Does your paper address subitem 1a-iii? \* Copy and paste relevant sections from manuscript title (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 Safe at Home Study: Protocol for a Randomized Controlled Trial of a Web-Based Decision Aid for Caregivers of Persons with Dementia with Firearm Access 1b) ABSTRACT: Structured summary of trial design, methods, results, and conclusions NPT extension: Description of experimental treatment, comparator, care providers, centers, and blinding status. 1b-i) Key features/functionalities/components of the intervention and comparator in the METHODS section of the ABSTRACT Mention key features/functionalities/components of the intervention and comparator in the abstract. If possible, also mention theories and principles used for designing the site. Keep in mind the needs of systematic reviewers and indexers by including important synonyms. (Note: Only report in the abstract what the main paper is reporting. If this information is missing from the main body of text, consider adding it) subitem not at all important essential Does your paper address subitem 1b-i? \* 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 "Participants will be randomized to view SiD or a control website (at their own pace)" 1b-ii) Level of human involvement in the METHODS section of the ABSTRACT Clarify the level of human involvement in the abstract, e.g., use phrases like "fully automated" vs. "therapist/nurse/care provider/physician-assisted" (mention number and expertise of providers involved, if any). (Note: Only report in the abstract what the main paper is reporting. If this information is missing from the main body of text, consider adding it) 1 2 3 5 subitem not at all important essential

Does your paper address subitem 1b-ii?

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

Participants will be randomized to view SiD or a control website (at their own pace)

1b-iii) Open vs. closed, web-based (self-assessment) vs. face-to-face assessments in the METHODS section of the ABSTRACT

Mention how participants were recruited (online vs. offline), e.g., from an open access website or from a clinic or a closed online user group (closed usergroup trial), 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). Clearly say if outcomes were self-assessed through questionnaires (as common in web-based trials). Note: In traditional offline trials, an open trial (open-label trial) is a type of clinical trial in which both the researchers and participants know which treatment is being administered. To avoid confusion, use "blinded" or "unblinded" to indicated the level of blinding instead of "open", as "open" in web-based trials usually refers to "open access" (i.e. participants can self-enrol). (Note: Only report in the abstract what the main paper is reporting. If this information is missing from the main body of text, consider adding it)

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# Does your paper address subitem 1b-iii?

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

"The Safe at Home (S@H) study is a two-armed randomized controlled trial that will test the impact of the SiD decision aid on caregivers of community-dwelling adults with ADRD who have firearm access. S@H will recruit 500 ADRD caregivers (age ≥18, fluent in English or Spanish) through online/social media advertisements and through relevant organizations. Participants will be randomized to view SiD or a control website (at their own pace); all participants will complete online follow-up questionnaires at 2 weeks, 2 months, and 6 months. "

# 1b-iv) RESULTS section in abstract must contain use data Report number of participants enrolled/assessed in each group, the use/uptake of the intervention (e.g., attrition/adherence metrics, use over time, number of logins etc.), in addition to primary/secondary outcomes. (Note: Only report in the abstract what the main paper is reporting. If this information is missing from the main body of text, consider adding it) subitem not at all important essential Clear selection Does your paper address subitem 1b-iv? 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 No because this is a protocol paper without data 1b-v) CONCLUSIONS/DISCUSSION in abstract for negative trials Conclusions/Discussions in abstract for negative trials: Discuss the primary outcome - if the trial is negative (primary outcome not changed), and the intervention was not used, discuss whether negative results are attributable to lack of uptake and discuss reasons. (Note: Only report in the abstract what the main paper is reporting. If this information is missing from the main body of text, consider adding it) $\bigcirc$ $\circ$ essential subitem not at all important Clear selection 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

"The S@H study is the first randomized trial of a firearm safety decision aid for ADRD caregivers. Results from this study will inform how best to support caregivers in decision-making regarding firearm safety. Further, results may guide approaches for recruiting caregivers and for dissemination of resources."

INTRODUCTION		

2a) In INTRODUCTION: Scient	ific bac	kground	l and exp	olanatio	n of ratio	onale
2a-i) Problem and the type of Describe the problem and the type as stand-alone intervention vs. in particular patient population? Go other interventions, replace or co intervention are provided in "Met	pe of sys ncorpora pals of th ompleme	stem/sol ated in br ne interve ent other	ution tha oader he ention, e.	alth care g., being	program more cos	? Intended for a st-effective to
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Does your paper address sub	item 2a	-i? *				
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"There is an urgent need for effective caregivers in decision making an regarding home safety. Ideally, do the broad, diverse community of delay nursing home placement, a	d behavi evelopm informa	ior chang ent and c I caregive	je to act d lissemina ers – cou	on those ation of s Id help re	decisions uch inter	s, especially ventions – to
2a-ii) Scientific background, ra Scientific background, rationale: object of the study (be sure to di conditions/diagnoses, if appropi and what is the context for this s study performed, potential impa comparator.	What is iscuss that is in the interior in the interior is the interior in the interior is the interior in the	known a ne use of otivation study, fro	bout the similar s for the st om which	(type of) systems f udy, i.e. v stakeho	system to for other what are to lder view	that is the the reasons for point is the
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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

"Providing resources to caregivers regarding firearm safety is of particular importance due to increased risk of suicide and risk related to behavior changes associated with ADRD ...."
"Online resources have the potential to reach a broad range of caregivers"

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

"The primary goal of the Safe at Home study is to assess the efficacy of the SiD online decision aid with respect to ability of the decision aid to assist caregivers on decision quality and behaviors regarding firearm access and injury reduction in the home of the person with ADRD. The study hypothesis is that the decision aid will increase the preparation and self-efficacy of informal caregivers to make and implement decisions that effectively address firearm access, thereby reducing firearm injury risk."

## **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

The S@H study is a two-armed randomized controlled trial.

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

no significant changes to trial thus far, not relevant to methods paper

# 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].

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#### 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

not relevant, study protocol for study with website that is intended to remain static

# 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

"To be eligible for participation in the study, participants must be ages 18 years or older, live in the United States, be fluent in English or Spanish, not in legal custody or institutionalized, and identify as informal caregivers for a community-dwelling person with ADRD who has firearm access. "Community-dwelling" is defined as a person with ADRD living in a private home and not in any type of care-giving facility (Figure 2). Participants must have internet access but do not need to live in the same home (or state) as the person with ADRD."

4a-i) Computer / Internet literacy is of explicitly clarified.	-	mplicit "d	de facto"	eligibility	criterion	- this should be
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Does your paper address sub Copy and paste relevant section "like this" to indicate direct quot providing additional information applicable/relevant for your stud "Participants must have internet	s from t es from not in tl ly	he manu your ma ne ms, or	nuscript)	, or elabo	orate on t	his item by
4a-ii) Open vs. closed, web-based vs. recruited (online vs. offline), e.g. if this was a purely web-based trintervention or for assessment), participant. In online-only trials, having multiple identities was pocookies, email confirmation, pho	face-to-f , from and ial, or the i.e., to veclarify if cossible cossible co	face asse n open ac ere were what degr participa or whethe	essments coess we face-to-f ree got th ants were er technic	s: Mention bsite or face com ne study face quasi-a cal or log	n how pa from a cli ponents team to k nonymou istical me	nic, and clarify (as part of the know the as and whether easures (e.g.,
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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

"The S@H study aims to enroll 500 English- and Spanish-speaking caregivers through social media recruitment (including Facebook, Instagram, and Google Ads) and relevant organizations in dementia care/aging, caregiver support, and firearms. "

"Potential participants who are eligible based on the online form provide contact information, and study staff then contact them by phone to validate identity (to reduce fraud or bot enrollment) and review eligibility and informed consent."

"Participants complete an online questionnaire before viewing the SiD decision aid or the control website (Figure 2, Table 1). After reviewing SiD or control, participants in both arms complete a second online questionnaire with questions about preparation for decision making, self-efficacy, and knowledge (Table 1). One week after the baseline session, participants receive a text or email reminder (depending on preference) with a hyperlink to the website to which they were randomized so they may visit it again if desired. Subsequent follow-up (at two weeks, two months, and six months) occurs via online questionnaires that are sent to participicants."

#### 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.

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# Does your paper address subitem 4a-iii?

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

"and review eligibility and informed consent (Multimedia Appendix 4)."

4b) Settings and locations where the data were collected

Does your paper address CONSORT subitem 4b? \*

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 can complete the study procedures at a time and in a place of their choosing, on a computer or other web-enabled device of their choosing"

4b-i) Report if outcomes were (self-)assessed through online questionnaires
Clearly report if outcomes were (self-)assessed through online questionnaires (as common in web-based trials) or otherwise.

1 2 3 4 5

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Clear selection

Does your paper address subitem 4b-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 complete an online questionnaire before viewing the SiD decision aid or the control website (Figure 2, Table 1). After reviewing SiD or control, participants in both arms complete a second online questionnaire with questions about preparation for decision making, self-efficacy, and knowledge (Table 1). One week after the baseline session, participants receive a text or email reminder (depending on preference) with a hyperlink to the website to which they were randomized so they may visit it again if desired. Subsequent follow-up (at two weeks, two months, and six months) occurs via online questionnaires that are sent to participants."

# 4b-ii) Report how institutional affiliations are displayed

Report how institutional affiliations are displayed to potential participants [on ehealth media], as affiliations with prestigious hospitals or universities may affect volunteer rates, use, and reactions with regards to an intervention. (Not a required item – describe only if this may bias results)

1 2 3 4 5 subitem not at all important 

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Does your paper address sub	item 4b	-ii?									
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"All advertisements contain similar messages and will direct individuals to a landing webpage with a description of the study, including the lead organization (University of Colorado)."											
5) The interventions for each including how and when they	-				ıllow rep	lication,					
5-i) Mention names, credential, affiliations of the developers, sponsors, and owners Mention names, credential, affiliations of the developers, sponsors, and owners [6] (if authors/evaluators are owners or developer of the software, this needs to be declared in a "Conflict of interest" section or mentioned elsewhere in the manuscript).											
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5-ii) Describe the history/developme evaluations (e.g., focus groups,	at no cos elopmer nt proce usability	nt proces ss of the testing),	ss applicati as these	on and p	revious f	ormative					
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## 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

"To address caregiver needs related to firearm access, and guided by theory [33-35], our team previously developed the online Safety in Dementia (SiD) online decision aid [36], which is available for public use at no cost. This online tool adheres to the International Patient Decision Aid Standards [35] and recommendations for user-friendly web design for older adults [37-39]. SiD is based on behavior change and decision-making theories [33,34] (Figure 1) and includes content to help caregivers understand options (with a balanced presentation of benefits and drawbacks) and then be motivated to take the action that works best for their situation. SiD provides parallel sections regarding driving safety and general home hazards to broaden reach and contextualize firearm safety. The SiD tool was translated into Spanish using a rigorous translation process including independent backtranslation to English by certified translators with adjudication of any discrepancies [40,41]. The pilot trial of SiD, in a national convenience sample of caregivers of community-dwelling people with ADRD, found 44% of caregivers reported safety concerns regarding firearm access and approximately 30% were currently considering options for what to do about firearm access [42]. SiD was developed to help caregivers work through these considerations - and this trial will test its efficacy in doing so. "

## 5-iii) Revisions and updating

Revisions and updating. Clearly mention the date and/or version number of the application/intervention (and comparator, if applicable) evaluated, or describe whether the intervention underwent major changes during the evaluation process, or whether the development and/or content was "frozen" during the trial. Describe dynamic components such as news feeds or changing content which may have an impact on the replicability of the intervention (for unexpected events see item 3b).

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# Does your paper address subitem 5-iii?

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

this is a study methods paper for the efficacy trial

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we did not include the source code for the website

5-vi) Digital preservation Digital preservation: Provide the change or disappear over the coarchived (Internet Archive, webc screenshots/videos alongside tarchived, consider creating dem	ourse of to citation.o the article	the years org, and/o e). As pag	; also ma or publish ges behir	ike sure t iing the s nd login s	the interv ource co screens c	rention is ade or cannot be
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5-vii) Access  Access: Describe how participal they had to pay (or were paid) or known, describe how participal ensure access for editors/review account or demo mode for review archiving purposes, see vi).	r not, wh its obtair wers/rea	ether the ned "acce ders, cor	y had to ess to the esider to	be a mer platform provide a	mber of s and Inte "backdo	pecific group. If ernet" [1]. To oor" login
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Does your paper address subitem 5-vii? \*

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 complete an online questionnaire before viewing the SiD decision aid or the control website (Figure 2, Table 1). After reviewing SiD or control, participants in both arms complete a second online questionnaire with questions about preparation for decision making, self-efficacy, and knowledge (Table 1). One week after the baseline session, participants receive a text or email reminder (depending on preference) with a hyperlink to the website to which they were randomized so they may visit it again if desired. Subsequent follow-up (at two weeks, two months, and six months) occurs via online questionnaires that are sent to participants. Participants receive incentives for completion of study measures (\$40 for completion of enrollment questionnaire, \$60 for 2-week and \$40 each for 2- and 6-month follow-up questionnaires)."

5-viii) Mode of delivery, features/functionalities/components of the intervention and comparator, and the theoretical framework

Describe mode of delivery, features/functionalities/components of the intervention and comparator, and the theoretical framework [6] used to design them (instructional strategy [1], behaviour change techniques, persuasive features, etc., see e.g., [7, 8] for terminology). This includes an in-depth description of the content (including where it is coming from and who developed it) [1]," whether [and how] it is tailored to individual circumstances and allows users to track their progress and receive feedback" [6]. This also includes a description of communication delivery channels and – if computer-mediated communication is a component – whether communication was synchronous or asynchronous [6]. It also includes information on presentation strategies [1], including page design principles, average amount of text on pages, presence of hyperlinks to other resources, etc. [1].

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## 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

"The study and its outcomes are based on the Ottawa Decision Support Framework (Figure 1) [33], which draws upon concepts from psychology, decision conflict, social support, and self-efficacy. The framework posits that decisional needs (i.e., knowledge, conflict/uncertainty, and values) affect decision quality, with the highest quality decision being one that is both informed by and reflective of the individual's values. Decision aids enhance decision quality by addressing unmet decisional needs, by: identifying the decision to be made, providing a balanced description of the risks and benefits of various options, assisting in clarifying personal values, and activating the individual for decision making [35,45]. Decision aids improve communication and knowledge and decrease decisional conflict and regret [46]. Application of the Health Belief Model [34] further expands this theoretical framework for how decision aids encourage action."

# 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.

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#### Does your paper address subitem 5-ix?

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 complete the study procedures at a time and in a place of their choosing, on a computer or other web-enabled device of their choosing."

"After reviewing SiD or control for as long as they want, participants in both arms complete a second online questionnaire with questions about preparation for decision making, self-efficacy, and knowledge (Table 1). One week after the baseline session, participants receive a text or email reminder (depending on preference) with a hyperlink to the website to which they were randomized so they may visit it again if desired"

# 5-x) Clarify the level of human involvement Clarify the level of human involvement (care providers or health professionals, also technical assistance) in the e-intervention or as co-intervention (detail number and expertise of professionals involved, if any, as well as "type of assistance offered, the timing and frequency of the support, how it is initiated, and the medium by which the assistance is delivered". It may be necessary to distinguish between the level of human involvement required for the trial, and the level of human involvement required for a routine application outside of a RCT setting (discuss under item 21 – generalizability). 1 2 3 4 5 subitem not at all important Clear selection

## Does your paper address subitem 5-x?

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

no human involvement offered

#### 5-xi) Report any prompts/reminders used

Report any prompts/reminders used: Clarify if there were prompts (letters, emails, phone calls, SMS) to use the application, what triggered them, frequency etc. It may be necessary to distinguish between the level of prompts/reminders required for the trial, and the level of prompts/reminders for a routine application outside of a RCT setting (discuss under item 21 – generalizability).

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# Does your paper address subitem 5-xi? \*

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

"One week after the baseline session, participants receive a text or email reminder (depending on preference) with a hyperlink to the website to which they were randomized so they may visit it again if desired"

5-xii) Describe any co-interventions (incl. training/support)

Describe any co-interventions (incl. training/support): Clearly state any interventions that are provided in addition to the targeted eHealth intervention, as ehealth intervention may not be designed as stand-alone intervention. This includes training sessions and support [1]. It may be necessary to distinguish between the level of training required for the trial, and the level of training for a routine application outside of a RCT setting (discuss under item 21 – generalizability.

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Does your paper address subitem 5-xii? \*

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

no cointerventions provided

subitem not at all important

6a) Completely defined pre-specified primary and secondary outcome measures, including how and when they were assessed

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

"The primary outcome is SiD's effect on preparation for decision making, a core element of the Ottawa Decision Support Framework"

"The Preparation for Decision Making Scale assesses perception of how useful a decision aid is in preparing for subsequent decision-making. Scores range from 1-5, calculated from the average of 10 constructs (each ranging from 1, "strongly disagree" to 5, "strongly agree"). The scale has total test reliability of 0.944 [54]. We will measure decision self-efficacy (self-confidence or belief in one's capability to make decisions) using the Decision Self-Efficacy Scale [55], as decision aids typically increase self-efficacy [56]. Transformed scores range from 0 (extremely low) to 100 (extremely high self-efficacy). In our pilot trial, participants had a mean score of 77.6 (SD 17.2) after viewing the SiD firearm section [42] (Table 1)."

"Key secondary outcomes, also linked to our theoretical framework (Figure 1), are in Table 1. Firearm access (and actions to reduce access) for the person with ADRD will be assessed with multi-point scales we have used previously [57,58], with binary categorization (any access to ≥1 firearm versus no access to any firearms) and progression towards reduced access (i.e., locking of additional firearm). This allows identification of smaller, albeit important, changes. We will assess firearm injury or near-injury in the person with ADRD or others in their home through questions about recent experiences (including incidents of threats with a firearm or "near-misses") and perceived risks. We will measure caregiving experience, both burden and positive experiences. Caregiver burden will be measured by the short-form (3-item) Zarit Burden Interview [59−61]. Benefits of caregiving will be measured by the Positive Aspects of Caregiving Scale [62−64]; scores range from 0-36, calculated from the sum of 9 items (measured on a 5-point scale ranging from 0, "disagree a lot" to 4, "agree a lot"), with higher scores indicating positive "

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].

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Does your paper address subitem 6a-i?

Copy and paste relevant sections from manuscript text

used pre-existing validated scales - see section above with text on methods

6a-ii) Describe whether and he defined/measured/monitored Describe whether and how "use" defined/measured/monitored (lo important process outcomes that	(includi	ng intens	sity of us ysis, etc.	e/dosage ). Use/ac	e) was doption m	
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6a-iii) Describe whether, how, obtained Describe whether, how, and whe (e.g., through emails, feedback f	n qualita	ative feed	back fro	m partici		•
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Does your paper address CONSORT subitem 6b? \*

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

NA trial underway, no changes yet

7a) How sample size was determined

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

7a-i) Describe whether and how expected attrition was taken into account when calculating the sample size

Describe whether and how expected attrition was taken into account when calculating the sample size.

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#### Does your paper address subitem 7a-i?

Copy and paste relevant sections from manuscript title (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 target sample size for this study (n=500), which includes recruitment via social media (n=250) and recruitment via relevant organizations (n=250), has been powered conservatively for the primary outcome to allow comparisons by recruitment groups and other participant characteristics. The large sample size will also protect against uncertainty in the estimate of the effect size given the small sample size (n=15) in the pilot study.[42] In the pilot trial of 15 caregivers, the mean Preparation for Decision Making scores (using 9 constructs) was 3.9 (SD 0.7) after SiD versus 3.6 (SD=0.2) in the control group; this corresponds to a standardized effect size of 0.42 (3.9 versus 3.6 with a pooled SD of 0.7; d=0.3/0.7=0.42) [42,76]. Based on t-test, with the proposed sample size of n=500 participants, analysis will be able to detect a d=0.42 effect size with very high power of >99% (alpha=0.05) between SiD and control for the primary outcome (preparation for decision making). Even if the effect is smaller (i.e., d=0.30) and the sample size is 80% (400 total) the power remains high at 84.9%. With a sample size of 500, there will be 125 in each of the final four analytic groups (stratified by recruitment group A versus B, SiD versus control). This sample will yield 80% power to detect any subgroup-specific standardized effect of d=0.36 or greater between SiD and control, including by recruitment group and other participant characteristics (using t-test and assuming sample sizes of 125 in each subgroup). Power calculations were computed using R version 4.0.2 [77]. "

7b) When applicable, explanation of any interim analyses and stopping guidelines

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

NA no stopping guidelines or interim analyses

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

"Enrolled participants are randomly assigned in equal numbers to the SiD intervention or control arm (Figure 2). Randomization is stratified between recruitment group A (social media/internet) and B (relevant organizations). To reduce bias and aim for balance across arms, participants are randomized into blocks of undisclosed size. The team biostatistician conceals allocation using a centralized, computer-generated list in REDCap [50]. The study team PI and one of the biostatisticians will be blinded; other members of the study team are unblinded."

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

"Enrolled participants are randomly assigned in equal numbers to the SiD intervention or control arm (Figure 2). Randomization is stratified between recruitment group A (social media/internet) and B (relevant organizations). To reduce bias and aim for balance across arms, participants are randomized into blocks of undisclosed size. The team biostatistician conceals allocation using a centralized, computer-generated list in REDCap [50]. The study team PI and one of the biostatisticians will be blinded; other members of the study team are unblinded."

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

Does your paper address CONSORT subitem 9? \*

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

"Enrolled participants are randomly assigned in equal numbers to the SiD intervention or control arm (Figure 2). Randomization is stratified between recruitment group A (social media/internet) and B (relevant organizations). To reduce bias and aim for balance across arms, participants are randomized into blocks of undisclosed size. The team biostatistician conceals allocation using a centralized, computer-generated list in REDCap [50]. The study team PI and one of the biostatisticians will be blinded; other members of the study team are unblinded."

10) Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventions

Does your paper address CONSORT subitem 10? \*

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

"Enrolled participants are randomly assigned in equal numbers to the SiD intervention or control arm (Figure 2). Randomization is stratified between recruitment group A (social media/internet) and B (relevant organizations). To reduce bias and aim for balance across arms, participants are randomized into blocks of undisclosed size. The team biostatistician conceals allocation using a centralized, computer-generated list in REDCap [50]. The study team PI and one of the biostatisticians will be blinded; other members of the study team are unblinded."

11a) If done, who was blinded after assignment to interventions (for example, participants, care providers, those assessing outcomes) and how NPT: Whether or not administering co-interventions were blinded to group assignment

11a-i) Specify who was blinded Specify who was blinded, and which blind the participants [1, 3] (this blind outcome assessors, those interventions (if any).	no wasn should b	t. Usuall e clearly	y, in web- acknow	ledged), l	out it ma	y be possible to
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Does your paper address subice Copy and paste relevant section "like this" to indicate direct quot providing additional information applicable/relevant for your stude "Enrolled participants are randomic control arm (Figure 2). Randomize media/internet) and B (relevant of arms, participants are randomized conceals allocation using a centiteam PI and one of the biostatist unblinded."	s from the sest from not in the ly assign action is organizated into black frailized, controlled in the sest from the sest from the left from	ne manua your man ne ms, or ned in ed stratified tions). To locks of u	nuscript) briefly ex qual num between reduce bundisclose-generate	or elaborates, or ela	orate on t by the iter he SiD int hent grou aim for b The team REDCap [	his item by m is not  ervention or p A (social alance across biostatistician 50]. The study
11a-ii) Discuss e.g., whether p "intervention of interest" and v Informed consent procedures (4 e.g., whether participants knew which one was the "comparator"	which o a-ii) can which in	ne was <sup>.</sup> create b	the "con iases an	nparator d certain	" expectat	tions - discuss
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11b) If relevant, description of the similarity of interventions (this item is usually not relevant for ehealth trials as it refers to similarity of a placebo or sham intervention to a active medication/intervention)

Does your paper address CONSORT subitem 11b?\*

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

"In the intervention arm, participants are directed via hyperlink to the SiD decision aid. Participants may view the site in the format they prefer (i.e. phone, computer, tablet) and are recommended to view the site for at least 5-10 minutes. A unique feature of the intervention is the intention to guide the participant through the decision or promote person-centric consideration of values and preferences [51].

In the active control arm, participants will be directed via hyperlink to the NIA Home Safety Checklist for Alzheimer's Disease website [47]. Similar to the intervention arm, participants in the control arm may view the site in the format they prefer (i.e. phone, computer, tablet). The control (NIA website) represents typical care, as it is an easily accessible website that consistently appears in the top five results of a Google search for "home safety dementia" and provides basic education about home safety in the context of dementia [47]. The checklist provides limited guidance on firearms ("Lock up or remove these potentially dangerous items from the home: ... Guns and other weapons, scissors, knives, power tools, and machinery") [47] without information about specific locking or disposal options. Unlike the intervention (SiD decision aid), the NIA website does not guide the individual through the decision or promote person-centric consideration of values and preferences, making it an appropriate control [51]."

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

"Descriptive statistics will be computed for baseline caregiver characteristics (including key covariates, Table 1), reporting on differences between: (a) intervention arm, (b) follow-up status, and (c) recruitment modality. To test the primary hypothesis that SiD causes higher immediate decision quality compared to control, we will test for a difference in the adjusted mean Preparation for Decision Making score at baseline after viewing SiD or control using multiple linear regression (MLR), with adjustment for baseline characteristics (l.e., age, gender, caregiver burden, severity of ADRD) as potential precision variables. Other components (decision quality, decision self-efficacy and knowledge) will be analyzed similarly. To assess longitudinal endpoints, linear mixed models (LMM) will replace MLR to assess changes in self-efficacy and knowledge over time by treatment arm, allowing an interaction of treatment arm and time.

To test the secondary hypothesis that SiD increases the odds that action is taken to reduce firearm access, a binomial-family logit-link generalized LMM will be employed with subject-specific random effects to account for correlations across measurements from the same individual, and with fixed effects to adjust for baseline characteristics. The interaction between intervention and time will be tested to assess if the effect of SiD varies over time. We will utilize interaction terms to identify potential differential effects of treatment will be assessed according to age, whether the caregiver lives with the person with ADRD (versus not), and recruitment group. Exploratory heterogeneity of treatment effect will also be conducted by caregiver gender, urban versus rural residence, and language (English versus Spanish). Additionally, we will examine longitudinal change in caregiver burden and reported safety concerns, and how these might affect treatment effect. Both multiplicity-adjusted and unadjusted p-values will be presented for these comparisons. "

# 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]).

treated in the statistical analysis (a complete case analysis is strongly discouraged, and simple imputation techniques such as LOCF may also be problematic [4]).							
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Does your paper address subitem 12a-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

"Substantial efforts will be made to minimize missing data in data collection, and our models will adjust for covariates associated with drop-out to attenuate bias. If necessary, we will also employ methods such as multiple imputation, pattern mixture models, and/or sensitivity analyses, following intent-to-treat principles."

12b) Methods for additional analyses, such as subgroup analyses and adjusted analyses

Does your paper address CONSORT subitem 12b? \*

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

""Descriptive statistics will be computed for baseline caregiver characteristics (including key covariates, Table 1), reporting on differences between: (a) intervention arm, (b) follow-up status, and (c) recruitment modality. To test the primary hypothesis that SiD causes higher immediate decision quality compared to control, we will test for a difference in the adjusted mean Preparation for Decision Making score at baseline after viewing SiD or control using multiple linear regression (MLR), with adjustment for baseline characteristics (l.e., age, gender, caregiver burden, severity of ADRD) as potential precision variables. Other components (decision quality, decision self-efficacy and knowledge) will be analyzed similarly. To assess longitudinal endpoints, linear mixed models (LMM) will replace MLR to assess changes in self-efficacy and knowledge over time by treatment arm, allowing an interaction of treatment arm and time.

To test the secondary hypothesis that SiD increases the odds that action is taken to reduce firearm access, a binomial-family logit-link generalized LMM will be employed with subject-specific random effects to account for correlations across measurements from the same individual, and with fixed effects to adjust for baseline characteristics. The interaction between intervention and time will be tested to assess if the effect of SiD varies over time. We will utilize interaction terms to identify potential differential effects of treatment will be assessed according to age, whether the caregiver lives with the person with ADRD (versus not), and recruitment group. Exploratory heterogeneity of treatment effect will also be conducted by caregiver gender, urban versus rural residence, and language (English versus Spanish). Additionally, we will examine longitudinal change in caregiver burden and reported safety concerns, and how these might affect treatment effect. Both multiplicity-adjusted and unadjusted p-values will be presented for these comparisons. ""

X26) REB/IRB Approval and Ethical Considerations [recommended as subheading under "Methods"] (not a CONSORT item)

X26-i) Comment on ethics co	mmitte	e approv	al			
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x26-ii) Outline informed conso Outline informed consent proce Checkbox, etc.?), and what infor be included in informed consent	dures e.o mation v	g., if cons was prov	ided (see			·
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X26-iii) Safety and security pr Safety and security procedures, the likelihood or detection of ha	incl. priv	acy cons				
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Does your paper address subitem X26-iii?

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 study team will work a Data Safety Monitoring Board, along with the IRB and the NIA, to monitor participant safety, evaluate study progress, review procedures for maintaining the confidentiality of data, and ensure the quality of data collection, management, and analyses. The following adverse and serious adverse events are monitored: any kind of negative interaction from intervention (e.g., verbal disagreement or physical fight), increased stress due to actions taken to reduce firearm access for person with dementia, physical injury sustained during actions to reduce firearm access for person with dementia, suicide ideation or attempt in caregiver or individual with dementia, hospitalization due to emotional factors (stress, depression, suicidal or homicidal ideation or intent), or hospitalization or surgery for physical injury sustained during actions to reduce firearm access for person with dementia. Each adverse event is graded by severity and relationship to intervention."

#### **RESULTS**

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

NA no results yet - methods paper

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

NA no results yet - methods paper

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.

subitem not at all important O O O essential

# 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

NA no results yet - methods paper

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

NA no results yet - methods paper

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  $\circ$ 0 0 subitem not at all important 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 NA no results yet - methods paper 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 NA no results yet - methods paper 15) A table showing baseline demographic and clinical characteristics for each NPT: When applicable, a description of care providers (case volume, qualification, expertise, etc.) and centers (volume) in each group Does your paper address CONSORT subitem 15? \* 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 NA no results yet - methods paper

15-i) Report demographics as	ssociate	d with d	ligital div	/ide issu	ies	
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16-ii) Primary analysis should Primary analysis should be inter only "users", with the appropriate 18-i).	nt-to-trea	t, second	lary anal	•			
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Does your paper address subitem 17a-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 NA no results yet - methods paper 17b) For binary outcomes, presentation of both absolute and relative effect sizes is recommended Does your paper address CONSORT subitem 17b? \* 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 NA no results yet - methods paper 18) Results of any other analyses performed, including subgroup analyses and adjusted analyses, distinguishing pre-specified from exploratory Does your paper address CONSORT subitem 18? \* 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 NA no results yet - methods paper 18-i) Subgroup analysis of comparing only users A subgroup analysis of comparing only users is not uncommon in ehealth trials, but if done, it must be stressed that this is a self-selected sample and no longer an unbiased sample from a randomized trial (see 16-iii). subitem not at all important essential

Does your paper address subitem 18-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

NA no results yet - methods paper

19) All important harms or unintended effects in each group (for specific guidance see CONSORT for harms)

Does your paper address CONSORT subitem 19? \*

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

NA no results yet - methods paper

#### 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 OOOO essential

## Does your paper address subitem 19-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

NA no results yet - methods paper

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X27) Conflicts of Interest (not a CONSORT item)

X27-i) State the relation of the study team towards the system being evaluated In addition to the usual declaration of interests (financial or otherwise), also state the relation of the study team towards the system being evaluated, i.e., state if the authors/evaluators are distinct from or identical with the developers/sponsors of the intervention.

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"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 "Study authors developed the SiD web-based decision aid being evaluated in this study. SiD is available to the public at no cost, and study authors have no financial gain from its use. " About the CONSORT EHEALTH checklist As a result of using this checklist, did you make changes in your manuscript? \* yes, major changes yes, minor changes What were the most important changes you made as a result of using this checklist? Your answer How much time did you spend on going through the checklist INCLUDING making \* changes in your manuscript approximately 2 hours of review/changes As a result of using this checklist, do you think your manuscript has improved? \* yes Other:

Copy and paste relevant sections from the manuscript (include quotes in quotation marks

Does your paper address subitem X27-i?

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  yes
o no
Other:
Clear selection
Any other comments or questions on CONSORT EHEALTH
Your answer
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!

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