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

The CONSORT-EHEALTH checklist is intended for authors of randomized trials evaluating webbased 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): Evsenbach 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/

doi: 10.2196/jmir.1923 PMID: 22209829

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Your nam	e	*
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First Last

Gladys Block

Primary Affiliation (short), City, Country \*

University of Toronto, Toronto, Canada

NutritionQuest, Berkeley, (

Your e-mail address * abc@gmail.com
gblock@berkeley.edu
Title of your manuscript *
Provide the (draft) title of your manuscript.
Alive-PD fully automated web- and internet-based diabetes prevention program:  Design and randomized controlled trial protocol
_ colg., and landonized colling also process.
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
o not submitted yet - in late draft status, just before submission
submitted to a journal but not reviewed yet
<ul> <li>submitted to a journal and after receiving initial reviewer comments</li> </ul>
submitted to a journal and accepted, but not published yet
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)
Other: JMIR Research Protocols
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-dig
number at the end of the DOI, to be found at the bottom of each published article in JMIR)
no ms number (yet) / not (yet) submitted to / published in JMIR
Other:

## **TITLE AND ABSTRACT**

1a) ITTLE: 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")
• yes
Other:
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.
1 2 3 4 5
subitem not at all important \( \cap \) \( \cap \) \( \cap \) essential
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 Alive-PD Study is a randomized parallel-group controlled trial of the Alive-PD program vs. a 6-month-delayed control group among participants with prediabetes."  "Alive-PD is delivered via an individualized website and interactive e-mails, and is supplemented by a smartphone application and automated phone and print modules."
<b>1a-ii) Non-web-based components or important co-interventions in title</b> Mention non-web-based components or important co-interventions in title, if any (e.g., "with telephone support").
1 2 3 4 5
subitem not at all important O O essential

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
We include Interactive Voice Response telephone counseling. However, we believe it is not essential to include this fact in the title.  "Automated phone and print The content of these Alive-PD modules was developed by nutrition and physical activity professionals, through collaboration with Stanford and Brown University experts. With these technologies, individually-tailored print and phone messages are delivered automatically, by computer algorithm. They make use of the
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 O O O essential
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 " diabetes prevention program"
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)

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subitem not at all important	$\bigcirc$	$\bigcirc$	$\bigcirc$	$\bigcirc$	•	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

"The program involves weekly tailored goal-setting, a team system and gamification, and other opportunities for interaction. An accompanying smartphone app supports goal-setting and activity planning."

"Participants were randomized by computer algorithm to begin the program immediately or after a 6-month delay."

### 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)

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### 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

"...Alive-PD, a one-year fully automated intervention delivered by email and web."

## 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 webbased 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
"HbA1c, glucose, lipids, BMI, waist circumference and blood pressure will be assessed at baseline, 3, 6, 9 and 12 months. Randomization and delivery of the intervention are independent of clinic staff, and clinic staff have been masked to the treatment group. "
<b>1b-iv) RESULTS section in abstract must contain use data</b> 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 O O essential
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  This is to be submitted to JMIR Research Protocols, so results are not presented.
"Participants (n=340) had a mean (SD) as follows: age 55 (8.9)y; BMI 31.1 (4.3); male 68.5%. "
<b>1b-v) CONCLUSIONS/DISCUSSION in abstract for negative trials</b> 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)
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INTRODUCTION						
INTRODUCTION						
2a) In INTRODUCTIO rationale	N: So	cier	itifi	c ba	ackground and explanation of	
alone intervention vs. incorport population? Goals of the inte	e type o rated i	of sy in bro n, e.	stem bade g., b	/solu r hea eing	tion that is object of the study: intended as star lth care program? Intended for a particular pati more cost-effective to other interventions, repla the intervention are provided in "Methods" und	ient ice oi
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indicate direct quotes from your information not in the ms, or large lighty-six million U.S. adult million in 2010 (1). Over half to Type 2 diabetes (T2DM) (interventions aimed at produ	ions from the cing land the ci	om tonuscient one with a content of the content of the content of the content on	he m ript), ain w third th pr send y cha d to y can p	or e hy th l, have e-diage nge reach	betes will eventually progress effective behavioral n dietary and physical activity large numbers of individuals lice long-term behavioral	nis" to

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

prevent progression of pre-diabetes to diabetes. It is a stand-alone intervention,

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 appropriate),

	/poir					for and what is the context for this specific study, formed, potential impact of findings [2]. Briefly justify
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There is a fairly extensive d or technology-only program						man-only, technology-assisted tion or weight loss.
Does your paper address of Copy and paste relevant second indicate direct quotes from y	CON ction our	NSOI ns fro man	RT s m th uscr	ubit e ma ipt),	em : anus	jectives or hypotheses  2b? * script (include quotes in quotation marks "like this" to laborate on this item by providing additional ne item is not applicable/relevant for your study
"We hypothesize that those significant reduction in fasti	in the	he in luco: post	terve se ai -bas	entio nd H seline	n gr bA1 e, ar	oup will achieve statistically c as compared to the delayed- nd that the intervention group will

## **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 Alive-PD Study is a randomized parallel-group controlled trial of the Alive-PD program vs. a 6-month-delayed control group among participants with prediabetes."
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
"A number of subjects who were eligible by one measure were found to have quite low fasting glucose, and a further restriction, glucose >= XX, was added after XX had been enrolled."
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?

"The smartphone app was introduced after approximately one-fourth of subjects had begun the trial, at which time its availability was announced to all participants."	
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	ιο
"Participants are recruited from a primary care health center of the Palo Alto Medical Foundation (PAMF), a community-based multi-specialty group practice	
in California. Primary care patients ages 30-69 years old and BMI >=27 with pre- diabetes as defined by a physician diagnosis, and/or fasting glucose in the pre-	
diabetic range (100-125 mg/dL) and/or Hemoglobin A1c in the pre-diabetic range (5.7-6.4%) at the baseline visit. Exclusion criteria were broad and intended to minimize confounding, optimize participant safety and minimize potential loss to	
follow-up. Table X provides detailed inclusion/exclusion criteria. "	
4a-i) Computer / Internet literacy	
Computer / Internet literacy is often an implicit "de facto" eligibility criterion - this should be explicitly clarified.	
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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	
"Potentially eligible participants were asked if they accessed their email or web at least once per week."	

## 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

clarify if participants were qu	asi-	anor	nymo	ous a	and v	am to know the participant. In online-only trials, whether having multiple identities was possible or okies, email confirmation, phone calls) were used to
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indicate direct quotes from y information not in the ms, or "At the baseline clinic visit, s fasting glucose, total and HI were assessed through poir Vantage point of care analyzanalyzer (21) respectively).  "At 3, 6, 9 and 12 months the repeated. An adverse event the second size of the second siz	etion our brie brie brie color brie color brie color brie color brie color col	s from mann fly e ed in thole care (20)	om thuscr xxpla form estero e who and biom onna	ne mainth, in which which which which we do not be the control of	or einny the consideration of	erides, and LDL by calculation I samples (Siemens DCA
may also bias results.						
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indicate direct quotes from y information not in the ms, or "Study invitation letters were	ction our brie	s from an all fly earlied	m thuscrexpla	ie ma ipt), in wh	or elny th	script (include quotes in quotation marks "like this" to laborate on this item by providing additional se item is not applicable/relevant for your study eligible participants, with phone gibility assessment, and invite

web-based trial, or there were face-to-face components (as part of the intervention or for

## 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
"After the baseline clinic and the initial account set-up, randomization and delivery of the intervention was conducted electronically by the Alive-PD program, completely independent of clinic staff."
<b>4b-i) Report if outcomes were (self-)assessed through online questionnaires</b> Clearly report if outcomes were (self-)assessed through online questionnaires (as common in webbased trials) or otherwise.
1 2 3 4 5 subitem not at all important O O essential
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  Outomes were assessed by biometric measures, stated several times already.
<b>4b-ii) Report how institutional affiliations are displayed</b> 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 \( \cap \cap \cap \cap \cap \cap \cap \cap
Does your paper address subitem 4b-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

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i-i) Mention names, cre Mention names, credenti	al, affiliations of	the develo	pers, sponsors	, and owners [6]	(if
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Two papers are cited, which	h de	scrib	e the	e rar	ndon	nized trial o	of prior vers	sions.		
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<b>5-iii) Revisions and updat</b> Revisions and updating. Cl	early									
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5-iv) Quality assurance m	etho	nde								
Provide information on qua			ance	met	hod	s to ensure	e accuracy	and quality	of information	
provided [1], if applicable.										
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information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"Two weekly Health Notes health information."	, dev	elop	ed b	y Ce	rtifie	ed Diabetes Educators, provide
5-v) Ensure replicability l capture video, and/or pro						arce code, and/or providing screenshots/screen- the algorithms used
video, and/or providing flow	vchar	ts of	the	algo	rithn	e, and/or providing screenshots/screen-capture ns used. Replicability (i.e., other researchers should Imark of scientific reporting.
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disappear over the course webcitation.org, and/or pub	of the dishir s can	e yea ng th not b	ars; a e so	also i urce chiv	make cod ed, c	cation, but as the intervention is likely to change or e sure the intervention is archived (Internet Archive, le or screenshots/videos alongside the article). As consider creating demo pages which are accessible
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information not in the ms, or briefly explain why the item is not applicable/relevant for your study

no
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5-vii) Access
Access: Describe how participants accessed the application, in what setting/context, if they had to
pay (or were paid) or not, whether they had to be a member of specific group. If known, describe how participants obtained "access to the platform and Internet" [1]. To ensure access for
editors/reviewers/readers, consider to provide a "backdoor" login account or demo mode for
reviewers/readers to explore the application (also important for archiving purposes, see vi).
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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
"After leaving the clinic, participants were sent an introductory email from the automated Alive-PD system, with further information and questionnaires. These
and all further online interactions were done at a time and place of their
choosing."
5-viii) Mada of dolivory foatures/functionalities/companents of the intervention and
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,
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
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
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
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
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
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
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].
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].
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].

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

This is described in detail or	ver s	SEVE	ral n	anes	:	
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5-ix) Describe use parame	ters					
						and optimal timing for use). Clarify what instructions
					e.g.,	regarding timing, frequency, heaviness of use, if
any, or was the intervention	use	d ad	IIDITI	um.		
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subitem not at all important	$\bigcirc$				ullet	essential
Does your paper address	subi	item	5-ix	?		
					anus	cript (include quotes in quotation marks "like this" to
						aborate on this item by providing additional
						e item is not applicable/relevant for your study
						lect one eating habits goal and
						jested each week. However,
they may select any number						
						k to report on their success at
the previous week's goals.						·
5-x) Clarify the level of hui	man	inve	olve	men	t	
						ders or health professionals, also technical
						tion (detail number and expertise of professionals
						ered, the timing and frequency of the support, how it
involved, if any, as well as "f	VUC					ce is delivered". It may be necessary to distinguish
		vhich	the	assi		
is initiated, and the medium	by w					
is initiated, and the medium between the level of human	by w invo	lven	nent	requ	ired	for the trial, and the level of human involvement setting (discuss under item 21 – generalizability).
is initiated, and the medium between the level of human	by w invo ation	lven out	nent side	requ of a	ired RCT	for the trial, and the level of human involvement
is initiated, and the medium between the level of human	by w invo	lven	nent side	requ	ired	for the trial, and the level of human involvement
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is initiated, and the medium between the level of human required for a routine applica	by winvo	outs 2	nent side	requ of a	ired RCT 5	for the trial, and the level of human involvement setting (discuss under item 21 – generalizability).
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is initiated, and the medium between the level of human required for a routine application subitem not at all important.  Does your paper address	by winvo	outs 2	nent side 3	requof a	ired RCT 5	for the trial, and the level of human involvement setting (discuss under item 21 – generalizability).

"Alive-PD is a fully-automa	ated in	terv	entic	on, w	/ith r	no human	coach	ing or	advice	. "		
										11		
5-xi) Report any prompts	/remi	nde	rs us	sed								
Report any prompts/remind to use the application, wha between the level of promp routine application outside	ders u t trigg ots/ren	sed: erec nind	Cla the ers r	rify i m, fr equi	equ red	ency etc. I for the tria	t may I, and	be nee	cessary vel of p	y to d romp	istinguisl ts/remin	٦
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information not in the ms, of "Mid-week reminders about											or your st	udy
participant engaged, and t responded by mid-week b	he ov	erall	obje	ectiv	e sa	lient. Parti	cipant	s who	have n	not		
smartphone of their self-re encouraged again to comm	porte	d mo	otiva	tions	for	joining the	progr	am, a	nd are			
encodiaged again to com	THE TO	1-2 (	guai	5 101	uic	remainder	OI LITE	WCCK				
										- 11		
5-xii) Describe any co-int Describe any co-intervention							-	- anv i	ntervei	ntions	that are	nrovided
in addition to the targeted e	eHealt	th in	terve	entio	n, a	s ehealth i	nterve	ntion r	may no	t be o	designed	as
stand-alone intervention. T distinguish between the lev												
application outside of a RC			_							allilli	g ioi a io	dille
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"Automated phone	e and	brint
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The content of these Alive-PD modules was developed by nutrition and physical activity professionals, through collaboration with Stanford and Brown University experts. With these technologies, individually-tailored print and phone messages are delivered automatically, by computer algorithm. They make use of the participant's baseline responses to factors such as barriers and motivations for behavior change; participant real-time drop-out or adherence status; on-going participant reports on success at goals undertaken in the Alive-PD program.

## 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

#### "Assessment of outcomes

Physiologic outcomes – which are the primary outcomes — are assessed by anthropometric or biometric measures, as stated above. Other outcomes are assessed by self-administered questionnaire, or through automated capture of use metrics. These include factors such as number of weeks in which a goal was chosen, number of goals chosen, number of goals achieved, number of team support messages sent, number of quizzes answered. Changes in physical activity and eating habits are assessed by online modified Block questionnaire.

## 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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subitem not at all important	$\bigcirc$	$\bigcirc$	•	$\bigcirc$	$\bigcirc$	essential

#### Does your paper address subitem 6a-i?

Copy and paste relevant sections from manuscript text

The paper does not directly address validation of the diet and activity questionnaires, as they are primarily behavior-change tools.

## 6a-ii) Describe whether and how "use" (including intensity of use/dosage) was defined/measured/monitored

Describe whether and how "use" (including intensity of use/dosage) was defined/measured/monitored

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goa	l wa	s ch	oser	n, nu	These include factors such as mber of goals chosen, number ages sent, number of quizzes
whe viev	n qu vs, f	ualita ocus	ative gro	feed ups)	ative feedback from participants was obtained back from participants was obtained (e.g., through
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ion	s fro	m m	anu		t text nclude satisfaction with the
1 Qf	w, where	w, and when quiviews, for a construction with the construction wit	w, and when the qualitative of use in the suppose of the suppose o	w, and when q when qualitative views, focus group 1 2 3 4 1 2 3 4 1 2 3	w, and when qualitative feed views, focus groups).  1 2 3 4 5  ubitem 6a-iii? ions from manuscript

## 6b) Any changes to trial outcomes after the trial commenced, with reasons

## Does your paper address CONSORT subitem 6b? \*

"Sample Size Sample size was estimated using data from Davis et al. (22), who conducted a dietary intervention on diabetics. They reported standard deviations (SD) for change in A1c of 0.9 to 1.4% for one arm of the study. With a SD of 1.4, a final sample of 268 would provide 80% power to detect a minimum detectable difference in change for A1c of 0.48. We planned to enroll 314 persons to achieve a completed sample of 268 after a 15% dropout rate."	
7a) How sample size was determined	
NPT: When applicable, details of whether and how the clustering by care provides o addressed	or centers was
7a-i) Describe whether and how expected attrition was taken into account whe the sample size  Describe whether and how expected attrition was taken into account when calculating size.  1 2 3 4 5	_
subitem not at all important O O essential	
Does your paper address subitem 7a-i? Copy and paste relevant sections from manuscript title (include quotes in quotation is to indicate direct quotes from your manuscript), or elaborate on this item by providing information not in the ms, or briefly explain why the item is not applicable/relevant for See above.	g additional

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

### Does your paper address CONSORT subitem 7b? \*

no	
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 mark indicate direct quotes from your manuscript), or elaborate on this item by providing addit information not in the ms, or briefly explain why the item is not applicable/relevant for you no	ional
8b) Type of randomisation; details of any restriction (such as b and block size)	locking
Does your paper address CONSORT subitem 8b? * Copy and paste relevant sections from the manuscript (include quotes in quotation mark indicate direct quotes from your manuscript), or elaborate on this item by providing addit information not in the ms, or briefly explain why the item is not applicable/relevant for you	ional
"Participants were randomized by computer algorithm, with blocked stratified randomization to achieve balance on sex, BMI (above/below 35) and race/ethnicity (white non-Hispanic, other)."	

# 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

"Participants were informed by the system about whether they had been randomized to begin the program immediately or after a 6-month delay. Those in the delayed group received no further contact until a reminder to complete a 3-month and 6-month follow-up questionnaire."

## 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

#### "Randomization and Blinding

After the baseline clinic and the initial account set-up, randomization and delivery of the intervention was conducted electronically by the Alive-PD program, completely independent of clinic staff. The web-based, online baseline questionnaire was required, to provide demographic and BMI information required for randomization, in addition to the capture of eating habits and physical activity data. Participants were randomized by computer algorithm, with blocked stratified randomization to achieve balance on sex, BMI (above/below

# 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, and who wasn't

Specify who was blinded, and who wasn't. Usually, in web-based trials it is not possible to blind the participants [1, 3] (this should be clearly acknowledged), but it may be possible to blind outcome assessors, those doing data analysis or those administering co-interventions (if any).

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						ne item is not applicable/relevant for your st
"Clinic staff obtaining the bi	ome	etric r	neas	sure	men	ts are blinded to treatment
group, while data analysts a						
11a-ii) Discuss e.g., wheth	er r	artio	cipa	nts k	(nev	wwhich intervention was the "intervention
interest" and which one w						
Informed consent procedure	es (4	a-ii)	can	crea	te b	ases and certain expectations - discuss e.g
	vhich	n inte	erver	ntion	was	the "intervention of interest" and which on
"comparator".						
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information not in the ms, or	brie	етіу е	expia	in w	ny tr	ne item is not applicable/relevant for your st
No. Treatment group was o	bvio	us.				

## 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? \*

No. Not similar.		
		,

## 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

"An intention-to-treat analysis is the primary approach taken on analyses of treatment effects on glycemic markers and weight. Variables missing at follow-up are assigned the last measured value. We also examine per-protocol analyses, in which effects are assessed among those providing follow-up biometric data and in relation to the participants' degree of interaction with the program."

#### 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]).



### Does your paper address subitem 12a-i? \*

"An intention-to-treat analysis is the primary approach taken on analyses of treatment effects on glycemic markers and weight. Variables missing at follow-up are assigned the last measured value. We also examine per-protocol analyses, in which effects are assessed among those providing follow-up biometric data and in relation to the participants' degree of interaction with the program."
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 see above
X26) REB/IRB Approval and Ethical Considerations [recommended as subheading under "Methods"] (not a CONSORT item)
X26-i) Comment on ethics committee approval  1 2 3 4 5

## Does your paper address subitem X26-i?

subitem not at all important O O O essential

	nu oi	the				by the Institutional Review edical Foundation (PAMF)."	
						<u>/</u>	
	oced was	lures	e.g	., if (	CO	sent was obtained offline or online (how? Chec 4a-ii). See [6] for some items to be included in	kbox,
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X26-iii) Safety and securit	y pro	oced	lure	s			
Safety and security procedu	ires,	incl.	priv	асу		asiderations, and any steps taken to reduce the	
Safety and security procedu	ires, rm (e	incl. e.g.,	priv edu	acy catio	on	nd training, availability of a hotline)	
Safety and security procedu	ires,	incl.	priv	асу	on		
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### **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

See	figure a				
					1

## 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) $^{\star}$

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

#### Intention to treat.

For some per protocol analyses, the following data are relevant: "Dropout rate (defined as failure to return for the in-clinic biometric measurements) as of the three-month clinic visit is 6% (20/340). Also as of the three-month time point, 78% of the Intervention group were still participating actively in the online program, by choosing or reporting on a goal, logging weight or activity, answering a quiz or sending a message to team members via the

#### 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.

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quotes in quotation marks "li this item by providing additio applicable/relevant for your s "Also as of the three-month	tion ke to onal study time onlin	s fro his" infor y e poi e pri ering	m th to ind mati nt, 7 ogra	dicat on n 8% o m, b	te di ot ir of th y ch	script or cite the figure number if applicable (include rect quotes from your manuscript), or elaborate on a the ms, or briefly explain why the item is not re Intervention group were still accosing or reporting on a goal, ending a message to team
14a) Dates defining t	the	pe	rio	ds d	of r	recruitment and follow-up
indicate direct quotes from y	ction our	s fro man	m th uscr	ie ma ipt),	anu: or e	14a? * script (include quotes in quotation marks "like this" to elaborate on this item by providing additional ne item is not applicable/relevant for your study
"Subjects were enrolled in the	ne s	tudy	betv	weer	n Fe	bruary and June, 2014."
	ents	s" fel	l into	the uter	stu hard	nto the study period dy period, e.g., significant changes in Internet dware or Internet delivery resources"
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	I
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
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information not in the ms, or briefly explain why the item is not applicable/relevant	for your study
INA	
15) A table showing baseline demographic and clinical cha	racteristics
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 CONSORT subitem 15? *	
Copy and paste relevant sections from the manuscript (include quotes in quotation indicate direct quotes from your manuscript), or elaborate on this item by providing	
information not in the ms, or briefly explain why the item is not applicable/relevant	
Yes, Table	
	I

## 15-i) Report demographics associated with digital divide issues

In ehealth trials it is particularly important to report demographics associated with digital divide issues,

such as age, education, ger participants, if known.	ider,	SOC	ial-e	conc	mic	status, computer/Internet/ehealth literacy of the
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each analysis and was 16-i) Report multiple "denominated Report multiple Report multiple "denominated Report multiple Repo	ominors" a	nato	r th	e a and ide d	naly prov lefini	cipants (denominator) included in ysis was by original assigned groups ride definitions tions: Report N's (and effect sizes) "across a range g., N exposed, N consented, N used more than x
times, N used more than y v	veek	s, N	part	icipa	ints '	'used" the intervention/comparator at specific pre- ative numbers per group). Always clearly define
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Primary analysis should be with the appropriate caveats						
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ndicate direct quotes from y	ction our	s fro man	m th uscr	ie ma ipt),	or el	cript (include quotes in quotation marks "like this" to aborate on this item by providing additional e item is not applicable/relevant for your study
						/2
17a) For each prima		_				
and the estimated e	-					ry outcome, results for each group, sprecision (such as 95% confidence
and the estimated entinterval)  Does your paper address Copy and paste relevant sendicate direct quotes from y	CON ction	it si ISOI s fro man	RT som thouser	ubite maipt),	em 'anus	s precision (such as 95% confidence  17a? * cript (include quotes in quotation marks "like this" to aborate on this item by providing additional
and the estimated edinterval)  Does your paper address Copy and paste relevant second control in the ms, or	CON ction our brie	ISOI s fro man	RT som the user xpla	ubite maipt), in wh	em 'anus	s precision (such as 95% confidence  17a? * cript (include quotes in quotation marks "like this" to
and the estimated entinterval)  Does your paper address Copy and paste relevant second control of the information not in the ms, or information not in the ms, or No. Results of the intervent in addition to primary/second metrics of use and intensity does not only refer to metric exposure metrics such as "a	CON ction /our briedion a	usoli since the	RT s m th uscr xpla ot pr com iical', lose, iion (sessi	ubitone maipt), in who reserves and the second seco	em ? anus or el ny th nted.	Tra?* cript (include quotes in quotation marks "like this" to aborate on this item by providing additional e item is not applicable/relevant for your study

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
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
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
<b>18-i)</b> 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).
1 2 3 4 5
subitem not at all important   O O O 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
40) All important harms or unintended offects in each group
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

19-i) Include privacy breaches, technical problems

participants, but also incider	nts s	uch	as p	ercei	ved	s does not only include physical "harm" to or real privacy breaches [1], technical problems, stended effects" also includes unintended positive
	1	2	3	4	5	
subitem not at all important	•	0	0	0	0	essential
indicate direct quotes from y	ction our	ns fro man	m th	e ma ipt),	or e	ccript (include quotes in quotation marks "like this" to aborate on this item by providing additional e item is not applicable/relevant for your study
Include qualitative feedback strengths and shortcomings	fron of the	n pa he ap	rticip oplica	ants ation	or c	pants or observations from staff/researchers abservations from staff/researchers, if available, on becially if they point to unintended/unexpected as for why people did or did not use the application
	1	2	3	4	5	
subitem not at all important	$oldsymbol{oldsymbol{\circ}}$	$\bigcirc$	$\bigcirc$	$\bigcirc$	$\bigcirc$	essential
indicate direct quotes from y	ction our	ns fro man	m th	e ma ipt),	or e	ccript (include quotes in quotation marks "like this" to aborate on this item by providing additional e item is not applicable/relevant for your study

## **DISCUSSION**

## 22) Interpretation consistent with results, balancing benefits and harms, and considering other relevant evidence

NPT: In addition, take into account the choice of the comparator, lack of or partial blinding, and unequal expertise of care providers or centers in each group

22-i) Restate study questions and process	oces d su	ss ou mma	u <b>tco</b> arize	mes	(us	e)			-	•	•
outcomes and process outc		•	,	4	_						
	1	2			5						
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NA		J.1.y C	7.1010		,		or applie			lo. you. o	tady
22-ii) Highlight unanswere	ad n	ew n	IIIES	tion	s ei	ınnest fiiti	ure rese	arch			
Highlight unanswered new			-					ui oii			
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indicate direct quotes from y information not in the ms, or											
NA		, ,	1 -		<i>y</i> -						<b>y</b>

## 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.

### 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

We will not have the duration or sample size to have a diabetes incidence endpoint. However, we do have the power to assess effectiveness for glycemic control and weight loss. Furthermore, the fact that the study is limited to subjects with email and internet access is a limitation. However, as of August 2011, 92% of American adults used email and the internet, including 93% of whites, 87% of African Americans and 88% of Hispanics, and over 50% had a smartphone. Finally, we recognize that wholly automated interactions, without person-to-

## 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

### Does your paper address subitem 21-i?

NA						
24 11 21				4.0		
21-II) Discuss if there were setting	ele	men	its ir	n the	RC	T that would be different in a routine application
prompts/reminders, more hu	ımar e ele	invo men	olvei	ment	t, tra	ould be different in a routine application setting (e.g., ining sessions or other co-interventions) and what e on use, adoption, or outcomes if the intervention is
	1		3	4	5	
subitem not at all important	•	0	$\bigcirc$	$\bigcirc$		essential
Does your paper address so					anus	script (include quotes in quotation marks "like this" to
						laborate on this item by providing additional ne item is not applicable/relevant for your study
NA		<i>y</i> -	1		<i>y</i> -	
OTHER INFORMATION						
23) Registration number and name of trial registry						

## Does your paper address CONSORT subitem 23? \*

yes	
24) Where the full trial protocol can be accessed, if available	
Does your paper address CONSORT subitem 24? * Cite a Multimedia Appendix, other reference, or copy and paste relevant sections from the (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript elaborate on this item by providing additional information not in the ms, or briefly explain w is not applicable/relevant for your study	t), or
NA.	
25) Sources of funding and other support (such as supply of dru	gs), role
of funders	
Does your paper address CONSORT subitem 25? * Copy and paste relevant sections from the manuscript (include quotes in quotation marks 'indicate direct quotes from your manuscript), or elaborate on this item by providing addition information not in the ms, or briefly explain why the item is not applicable/relevant for your	nal
"Funding. Research reported in this publication was supported by the National Institute of Nursing Research of the National Institutes of Health under Award Number R44NR012617."	

<b>X27-i)</b> 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.						
	1	2	3	4	5	
subitem not at all important	0	0	0	0	•	essential
indicate direct quotes from ye	tion our	s fro man	m th	ie ma ipt),	or el	ccript (include quotes in quotation marks "like this" to aborate on this item by providing additional e item is not applicable/relevant for your study
	PD p	rogr	am.	Rob	ert R	co-owners of NutritionQuest, comanelli, Kristen Azar, Heather of interest."
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