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

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

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

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

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

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

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

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

Mandatory reporting items are marked with a red *.

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

YOUR ANSWERS WILL BE PUBLISHED AS A SUPPLEMENTARY FILE TO YOUR PUBLICATION IN JMIR AND ARE CONSIDERED PART OF YOUR PUBLICATION (IF ACCEPTED).

Please fill in these questions diligently. Information will not be copyedited, so please use proper spelling and grammar, use correct capitalization, and avoid abbreviations.

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

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

Eysenbach G, CONSORT-EHEALTH Group

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

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

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

* Required	
•	
Your name *	
First Last	
Jinshuo Li	
Primary Affiliation (short), C	City, Country *
University of Toronto, Toronto,	Canada
University of York, York,	
Your e-mail address *	
abc@gmail.com	
jinshuo.li@york.ac.uk	
Title of your manuscript *	
Provide the (draft) title of your	manuscript.
Type 2 Diabetes: a randomis	ed-control trial
Type 2 Diabetes: a randomis	ed-control trial
Article Preparation Status/S	
Article Preparation Status/S At which stage in your article p	Stage * oreparation are you currently (at the time you fill in this form)
Article Preparation Status/S At which stage in your article p ○ not submitted yet - in early	Stage * oreparation are you currently (at the time you fill in this form)
Article Preparation Status/S At which stage in your article p onot submitted yet - in early not submitted yet - in late d	Stage * preparation are you currently (at the time you fill in this form) draft status traft status, just before submission
Article Preparation Status/S At which stage in your article p onot submitted yet - in early not submitted yet - in late d submitted to a journal but r	Stage * preparation are you currently (at the time you fill in this form) draft status traft status, just before submission
Article Preparation Status/S At which stage in your article p onot submitted yet - in early not submitted yet - in late d submitted to a journal but r	Stage * preparation are you currently (at the time you fill in this form) draft status lraft status, just before submission not reviewed yet
Article Preparation Status/S At which stage in your article p onot submitted yet - in early not submitted yet - in late d submitted to a journal but r	Stage * preparation are you currently (at the time you fill in this form) draft status fraft status, just before submission not reviewed yet after receiving initial reviewer comments
Article Preparation Status/S At which stage in your article p onot submitted yet - in early not submitted yet - in late d submitted to a journal but r submitted to a journal and a	Stage * preparation are you currently (at the time you fill in this form) draft status fraft status, just before submission not reviewed yet after receiving initial reviewer comments
Article Preparation Status/S At which stage in your article p onot submitted yet - in early not submitted yet - in late d submitted to a journal but r submitted to a journal and a submitted to a journal and a	Stage * preparation are you currently (at the time you fill in this form) draft status fraft status, just before submission not reviewed yet after receiving initial reviewer comments
Article Preparation Status/S At which stage in your article p onot submitted yet - in early not submitted yet - in late d submitted to a journal but r submitted to a journal and a submitted to a journal and a published Other:	Stage * preparation are you currently (at the time you fill in this form) draft status fraft status, just before submission not reviewed yet after receiving initial reviewer comments
Article Preparation Status/S At which stage in your article p onot submitted yet - in early onot submitted yet - in late d submitted to a journal but r submitted to a journal and a submitted to a journal and a published Other: Journal * If you already know where you	Stage * preparation are you currently (at the time you fill in this form) draft status fraft status, just before submission not reviewed yet after receiving initial reviewer comments

Other:	
L	
If this is a tracking nuathor in J	pt tracking number * JMIR submission, please provide the manuscript tracking number under "other" (The ms umber can be found in the submission acknowledgement email, or when you login as MIR. If the paper is already published in JMIR, then the ms tracking number is the fourer 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) TI	TLE: Identification as a randomized trial in the
	TEE. Identification as a randomized that in the
title	
I.e does th	your paper address CONSORT item 1a? * e title contain the phrase "Randomized Controlled Trial"? (if not, explain the reason unde
•	
I.e does th "other")	
I.e does th "other") • yes	
I.e does th "other") • yes Other:	
I.e does the "other") yes Other: 1a-i) Iden Identify the in the title. Intervention "electronic worlds). Us product na	e title contain the phrase "Randomized Controlled Trial"? (if not, explain the reason unde
I.e does the "other") yes Other: 1a-i) Iden Identify the in the title. Intervention "electronic worlds). Us product na	tify the mode of delivery in the title e mode of delivery. Preferably use "web-based" and/or "mobile" and/or "electronic game. Avoid ambiguous terms like "online", "virtual", "interactive". Use "Internet-based" only if in includes non-web-based Internet components (e.g. email), use "computer-based" or "only if offline products are used. Use "virtual" only in the context of "virtual reality" (3-Dase "online" only in the context of "online support groups". Complement or substitute times with broader terms for the class of products (such as "mobile" or "smart phone"
I.e does the "other") yes Other: 1a-i) Iden Identify the in the title. Intervention "electronic worlds). Use product nationstead of	tify the mode of delivery in the title e mode of delivery. Preferably use "web-based" and/or "mobile" and/or "electronic game. Avoid ambiguous terms like "online", "virtual", "interactive". Use "Internet-based" only if in includes non-web-based Internet components (e.g. email), use "computer-based" or only if offline products are used. Use "virtual" only in the context of "virtual reality" (3-Dase "online" only in the context of "online support groups". Complement or substitute times with broader terms for the class of products (such as "mobile" or "smart phone" "iphone"), especially if the application runs on different platforms.

"Cost-effectiveness of facilitated access to a self-management website (HeLP-Diabetes) compared to usual care for adults with Type 2 Diabetes: a randomised-control trial"	
Type 2 Diabetes, a fandomised-control thai	
a-ii) Non-web-based components or important co-interventions	
1ention non-web-based components or important co-interventions in telephone support").	itle, if any (e.g., "with
1 2 3 4 5	
ubitem not at all important 🔾 💿 🔾 🔾 essential	
oes your paper address subitem 1a-ii?	
opy and paste relevant sections from manuscript title (include quotes nis" to indicate direct quotes from your manuscript), or elaborate on the dditional information not in the ms, or briefly explain why the item is r	nis item by providing
our study "Cost-effectiveness of facilitated access to a self-management	
website (HeLP-Diabetes) compared to usual care for adults with	
Type 2 Diabetes: a randomised-control trial"	
a-iii) Primary condition or target group in the title lention primary condition or target group in the title, if any (e.g., "for c	hildren with Type I Diahetos"
xample: A Web-based and Mobile Intervention with Telephone Supportion in the title, if any (e.g., 101 c.) in the title, if any (e.g., 101 c.) is ample: A Web-based and Mobile Intervention with Telephone Supportion in the title, if any (e.g., 101 c.) is a supportion with Telephone Supportion in the title, if any (e.g., 101 c.) is a supportion with Telephone Supportion in the title, if any (e.g., 101 c.) is a supportion with Telephone Supportion with Telephone Supportion in the title, if any (e.g., 101 c.) is a supportion with Telephone Supportion in the title, if any (e.g., 101 c.) is a supportion with Telephone Supportion with Telephone Supportion in the title, if any (e.g., 101 c.) is a supportion with Telephone Supportion in the title, if any (e.g., 101 c.) is a supportion with Telephone Supportion in the title, if a support in the title, if a	
1 2 3 4 5	
ubitem not at all important \bigcirc \bigcirc \bigcirc essential	
oes your paper address subitem 1a-iii? *	

"Cost-effectiveness of facilitated access to a self-management website (HeLP-Diabetes) compared to usual care for adults with Type 2 Diabetes: a randomised-control trial"
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)
1 2 3 4 5
subitem not at all important \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 participants were then randomised to either usual care plus nurse-facilitated access to HeLP-Diabetes, an interactive, theoretically-informed web-based self-management programme, or to usual care plus access to a comparator web-site containing basic information only."
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 4 5 subitem not at all important O O • 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

"The participants were then randomised to either usual care plus nurse-facilitated access to HeLP-Diabetes, an interactive, theoretically-informed web-based self-management programme, or to usual care plus access to a comparator web-site containing basic information only."

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)

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

"Adults aged 18 or over with a diagnosis of type 2 diabetes and registered with the 21 participating general practices (primary care) in England, UK, were approached."

"Participants were blinded to their allocation. The participants' intervention costs and wider health care resource use were collected as well as two self-reported health-related quality of life measures: the Problem Areas in Diabetes (PAID) Scale and EQ-5D-3L. EQ-5D-3L was then used to calculate quality-adjusted life years (QALYs)."

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)

1 2 3 4 5

Copy and paste rele "like this" to indicat	e direct quotes from you	manuscript abstract (include ir manuscript), or elaborate of fly explain why the item is no	on this item by providing
intervention group The use data wer Because this is a factor that had dir	cost-effectiveness analy	group." iical effectiveness article. rsis, the use data wasn't a the running costs are fixed	
1b-v) CONCLUSIO	NS/DISCUSSION in al	hatraat for pogative trials	
negative (primary onegative results are abstract what the n	ssions in abstract for ne utcome not changed), a attributable to lack of u ain paper is reporting. I	gative trials: Discuss the prii nd the intervention was not i iptake and discuss reasons. f this information is missing	used, discuss whether (Note: Only report in the
negative (primary onegative results are abstract what the note consider adding it)	ssions in abstract for ne utcome not changed), a attributable to lack of u	gative trials: Discuss the prii nd the intervention was not i iptake and discuss reasons. f this information is missing	used, discuss whether (Note: Only report in the
negative (primary of negative results are abstract what the not consider adding it)	esions in abstract for ne utcome not changed), a attributable to lack of u aain paper is reporting. I 1 2 3 4	gative trials: Discuss the prind the intervention was not uptake and discuss reasons. If this information is missing	used, discuss whether (Note: Only report in the
negative (primary of negative results are abstract what the not consider adding it) subitem not at all in the properties of the propertie	ssions in abstract for ne atcome not changed), a attributable to lack of usin paper is reporting. In the properties of the state of the	gative trials: Discuss the prind the intervention was not uptake and discuss reasons. If this information is missing	used, discuss whether (Note: Only report in the from the main body of text) e quotes in quotation mark on this item by providing

INTRODUCTION

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

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

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

Does your paper address subitem 2a-i? *

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

"Patient education and self-management support has been identified as a priority for global health in recent years and has the potential to both improve outcomes and reduce costs. However, internationally, uptake of self-management education remains low, partly due to logistic problems with attending courses. Web-based self-management support has the potential to increase uptake by overcoming some of the logistic problems associated with other forms of delivery, as it can be accessed at home, at the

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

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

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

"However, internationally, uptake of self-management education remains low, partly due to logistic problems with attending courses." "Web-based self-management support has the potential to increase uptake by overcoming some of the logistic problems associated with other forms of delivery, as it can be accessed at home, at the user's convenience."

"unlike telephone-based or face-to-face education where labour costs account for a substantial proportion of total cost."

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 aim of this paper is to present the health economic analysis of this comparison based on the data collected in the trial and to examine the cost-effectiveness of facilitated access to HeLP-Diabetes from the perspective of the National Health Services (NHS) and personal social services (PSS), following the National Institute of Health and Care Excellence (NICE) guidance."

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 HeLP-Diabetes trial was a multi-centre, two-arm individually randomised controlled trial carried out in primary care settings in England, UK."

"Randomisation was conducted in a 1:1 ratio"

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

Does your paper address CONSORT subitem 3b? *

There was no changes to the	e methods a	fter the pr	rotocol was	agreed
and the start of the trail."				•

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 There was no such events. 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 "Participants were adults, aged 18 years or over, registered with participating practices, and diagnosed with T2DM. People, who were unable to provide informed consent (e.g. due to psychosis or cognitive impairment), or unable to use the intervention (e.g. due to physical, sensory or intellectual impairment, or inability to understand basic spoken or written English), or terminally lil, or currently participating in a trial of an alternative self-management intervention, were excluded." 4a-i) Computer / Internet literacy Computer / Internet literacy is often an implicit "de facto" eligibility criterion - this should be explicitly clarified.	Bug fixes, Downtimes, (description of changes intervention or compara	imes, Content Changes ontent Changes: ehealth systems are often dynamic systems. A o methods therefore also includes important changes made on the for during the trial (e.g., major bug fixes or changes in the functionality or "unexpected events" that may have influenced study design such as staff is/downtimes, etc. [2].
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 There was no such events. 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 "Participants were adults, aged 18 years or over, registered with participating practices, and diagnosed with T2DM. People, who were unable to provide informed consent (e.g. due to psychosis or cognitive impairment), or unable to use the intervention (e.g. due to physical, sensory or intellectual impairment, or inability to understand basic spoken or written English), or terminally ill, or currently participating in a trial of an alternative self-management intervention, were excluded." 4a-i) Computer / Internet literacy Computer / Internet literacy is often an implicit "de facto" eligibility criterion - this should be		1 2 3 4 5
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 There was no such events. 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 "Participants were adults, aged 18 years or over, registered with participating practices, and diagnosed with T2DM. People, who were unable to provide informed consent (e.g. due to psychosis or cognitive impairment), or unable to use the intervention (e.g. due to physical, sensory or intellectual impairment, or inability to understand basic spoken or written English), or terminally ill, or currently participating in a trial of an alternative self-management intervention, were excluded." 4a-i) Computer / Internet literacy Computer / Internet literacy is often an implicit "de facto" eligibility criterion - this should be	subitem not at all impo	ant O O • O essential
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 "Participants were adults, aged 18 years or over, registered with participating practices, and diagnosed with T2DM. People, who were unable to provide informed consent (e.g. due to psychosis or cognitive impairment), or unable to use the intervention (e.g. due to physical, sensory or intellectual impairment, or inability to understand basic spoken or written English), or terminally ill, or currently participating in a trial of an alternative self-management intervention, were excluded." 4a-i) Computer / Internet literacy Computer / Internet literacy is often an implicit "de facto" eligibility criterion - this should be	Copy and paste relevanto indicate direct quote information not in the n	sections from the manuscript (include quotes in quotation marks "like this" from your manuscript), or elaborate on this item by providing additional s, or briefly explain why the item is not applicable/relevant for your study
"Participants were adults, aged 18 years or over, registered with participating practices, and diagnosed with T2DM. People, who were unable to provide informed consent (e.g. due to psychosis or cognitive impairment), or unable to use the intervention (e.g. due to physical, sensory or intellectual impairment, or inability to understand basic spoken or written English), or terminally ill, or currently participating in a trial of an alternative self-management intervention, were excluded." 4a-i) Computer / Internet literacy Computer / Internet literacy is often an implicit "de facto" eligibility criterion - this should be	Does your paper addi Copy and paste relevan to indicate direct quote	ess CONSORT subitem 4a? * sections from the manuscript (include quotes in quotation marks "like this" from your manuscript), or elaborate on this item by providing additional
Computer / Internet literacy is often an implicit "de facto" eligibility criterion - this should be	"Participants were adeparticipating practices were unable to provid cognitive impairment) physical, sensory or in basic spoken or writted participating in a trial of the participating in a t	ts, aged 18 years or over, registered with and diagnosed with T2DM. People, who informed consent (e.g. due to psychosis or or unable to use the intervention (e.g. due to ellectual impairment, or inability to understand English), or terminally ill, or currently
1 2 3 4 5	Computer / Internet lite	acy is often an implicit "de facto" eligibility criterion - this should be
subitem not at all important \bigcirc \bigcirc \bigcirc essential	suhitem not at all imno	
- Constitution at all important C C C C C C C C C C C C C C C C C C C		ant (/ (/ (/ (/ (/ (/ / (/ / (/

information not in the ms,										ble	/rele	evant	for yo	our stu	dy
"There were no exclusion diabetes control, previou education, computer and internet at home."	s expe	erien	се	of se	elf-m	nanage	ment		el of						
4a-ii) Open vs. closed, w	veb-ba	ased	d vs	. fa	ce-t	o-face	e asso	essn	nents	s:					
Open vs. closed, web-base (online vs. offline), e.g., fro web-based trial, or there w assessment), i.e., to what clarify if participants were whether technical or logist to detect/prevent these.	om an o ere fac degree quasi-	oper ce-to e go -ano	n ac o-fac t the nym	cess ce co stu nous	s we omp idy t s and	ebsite o conents eam to d wheth	or fron s (as p knov ner ha	n a c part o v the oving	linic, a of the partion multi	and int cipa	l clai erve ant. l ider	rify if ntion In on ntities	this vor for for fine-or swas	vas a p r nly tria possik	urely s, le or
	1	2	3	4	5										
subitem not at all importar	nt ()	\bigcirc	•	\bigcirc											
Does your paper address			1 4a	-ii?	*										
Does your paper address Copy and paste relevant so to indicate direct quotes fr information not in the ms, "participants were recruit "The intervention consist Diabetes. Facilitation co with practice nurses. In to log on, set a user nam content of the website." At the time of the trial, the	ections or brief ed in the ed of forms noistee this apple and	s fro our n efly e the p facili d of opoir pas	n 4a om the nanu explai orac tate an i atme swo	-ii? he muscr ain v tices d ac ntro ent p rd, a	* nanu ipt), why s." cces duc patie	uscript or elab the iten s to He tory tra nts we introdu	(inclusionates in is not be the control of the cont	sessown lo the	his ite oplica ion how	em	by p	rovic	ling ad	ddition	al
Copy and paste relevant so to indicate direct quotes from information not in the ms, "participants were recruit "The intervention consist Diabetes. Facilitation cowith practice nurses. In to log on, set a user name content of the website." At the time of the trial, the therefore participants contents to the contents of the trial, the therefore participants contents to the contents of the trial, the therefore participants contents of the con	ection: rom yo or brie eed in t eed of f nsiste this ap ee and ee webs uld onl	s fro our n efly e the p acili d of poir pas site	n 4a om ti nanu orac tate an i ttme swo was	he muscr ain votices d acontro ent pord, a	* nanuipt), why s." cces ductoratie and public us	uscript or elak the iten s to He tory tra nts we introdu olicly ac ser nan	(inclusionates in is not be the control of the cont	sessown lo the	his ite oplica ion how	em	by p	rovic	ling ad	ddition	al
Copy and paste relevant so to indicate direct quotes fr information not in the ms, "participants were recruit "The intervention consist Diabetes. Facilitation co with practice nurses. In to log on, set a user nam content of the website." At the time of the trial, the	ections on your or brief ted in the ed of for this apple and the websuld only the ection of the ecti	s frour not fly of the process from the	n 4a om ti nanu prace prace tate an i ntme swo was t up ecr	he muscrain values tices date not pord, a not other to the control of the control	* nanu ipt), why s." cces duc patie and pub ir us	uscript or elab the iten s to He tory tra nts we introdu blicly ac ser nan t bw part he info	(inclusionates in is not be the control of the cont	sessown loo the	ion how lp	em ble	by p /rele	rovice evant or rec ntati	ling ad for yo cruitm on as	ent an appen	al dy d in dix,
Copy and paste relevant so to indicate direct quotes fr information not in the ms, "participants were recruit "The intervention consist Diabetes. Facilitation co with practice nurses. In to log on, set a user nam content of the website." At the time of the trial, the therefore participants contents of the trial, the therefore participants of the trial, the trial the	ections on your or brief ted in the ed of formation of the education of th	s frour not fly of the process from the	n 4a pm ti nanu practate an i ntme swo was t up ecr	he muscrain values described acountrol the motor than the control than the	* nanuipt), why s." cces duct patie and pub ir us	uscript or elab the iten s to He tory tra nts we introdu blicly ac ser nan t bw part he info	(inclusionates in is not be the control of the cont	sessown loo the	ion how lp	em ble	by p /rele	rovice evant or rec ntati	ling ad for yo cruitm on as	ent an appen	al dy d in dix,

Does your paper address subitem 4a-iii?

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

"Both these self-reported outcome measures were collected online" "Health care resource use...was collected for both groups in three sections using pre-designed questionnaires. The majority of information pertaining to service use and participants' prescriptions were extracted from participants' medical records by practice or research nurses. The remainder of the service use data were collected retrospectively from participants using a self-report questionnaire online."

4b-i) Report if outcomes were (self-)assessed through online questionnaires

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	0	0	•	0	0	essentia

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

"Both these self-reported outcome measures were collected online at baseline, three months, and 12 months follow-up."

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)

							_				
subitem not at all	important O	•	0	0	0	essentia	nl 				
Does your paper											
Copy and paste re to indicate direct nformation not in	quotes from yo	our n	nanı	ıscr	ipt),	or elabor	ate on	this ite	n by provi	iding additio	nal
UCL's logo was	displayed on p	atie	nt in	forn	natio	on sheet.	NHS			,	,
Whittington's na	me was displa	yed (on th	ne ir	nter	vention w	ebsite.				
•											า
details to they were 5-i) Mention nar Mention names, cauthors/evaluator	allow reactually actually mes, credential, affilings are owners of	pli / a al, a ation	ica dr ffilians of	nii nii atio f the	on nis	of the developers, he softwa	velope sponso	rs, spo	omsors, ar	nd when	
details to they were 5-i) Mention nar Mention names, c authors/evaluator	allow reactually actually mes, credential, affilings are owners of	pli / a al, a ation	ica dr ffilians of	nii nii atio f the	on nis	of the developers, he softwa	velope sponso	rs, spo	omsors, ar	nd when	
details to they were 5-i) Mention nar Mention names, c authors/evaluator	allow reactually nes, credential, affiliars are owners on or mentioned	pli / a al, a ation	dr ffilians of evelouewh	ationil ationil f theoper ere	ons on the of the one	of the developers, he softwa	velope sponso	rs, spo	omsors, ar	nd when	
details to they were 5-i) Mention nar Mention names, c authors/evaluator of interest" sectio	allow reactually mes, credential, affilial are owners on or mentioned	pli / a al, a ation or de d els 2	dr ffilians of evelo ewh	nii nii atio f the per ere	ons on the office of the office of the office of the office office of the office offic	of the developers, he software manus	velope sponsoare, this cript).	rs, spo	omsors, ar	nd when	
details to they were 5-i) Mention nar Mention names, c authors/evaluator of interest" sectio	allow reactually mes, credential, affilial are owners on or mentioned	pli / a al, a ation or de d els 2	dr ffilians of evelo ewh	nii nii atio f the per ere	ons on the office of the office of the office of the office office of the office offic	of the developers, he software manus	velope sponsoare, this cript).	rs, spo	omsors, ar	nd when	
details to they were 5-i) Mention nar Mention names, c authors/evaluator of interest" section	allow reactually mes, credential, affiliates are owners on or mentioned 1	pli/ a al, a ation de dels	dr ffilians of velo ewh	niii niii atio f the per ere 4	ons on the office of the office of the office of the office office of the office offic	of the developers, he software manus	velope sponsoare, this cript).	rs, spo	omsors, ar	nd when	
5) The int details to they were 5-i) Mention names, of authors/evaluator of interest" section subitem not at all Does your paper Copy and paste re	allow reactually mes, credential, affiliates are owners on or mentioned important or address subjects.	al, a ation or de dels	ffilians of velocewhas 3	niii atio f the per ere 4 •	on nis	of the developers, he softwahe manus	velope sponso are, this cript).	rs, spoors, and needs	onsors, and owners [6 to be dec	nd when nd owners is (if lared in a "C	onflic
details to they were 5-i) Mention nar Mention names, of authors/evaluator of interest" section	allow reactually mes, credential, affiliates are owners on or mentioned important of address subselevant section quotes from your actually	al, a ation or ded dels	dr ffilians of veloewh 3	nil atio f the per ere 4 ? ne ne nuscr	on nis	of the developers, he softwathe manus	velope sponso are, this cript).	rs, spoors, and needs	onsors, are owners [6 to be dec	nd when a downers on marks "like iding addition and when a difference in a dif	onflic e this nal

5-ii) Describe the history/development process

Here 'we' meant our research team. The website was the first

"EM is the Managing Director of a not-for-profit Community Interest Company established to disseminate HeLP-Diabetes across the

Diabetes, or HeLP-Diabetes."

phase of a large project.

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

Copy and paste rele	ddress subitem 5-ii? vant sections from the manuscript (include quotes in quotation marks "like this"
	otes from your manuscript), or elaborate on this item by providing additional ems, or briefly explain why the item is not applicable/relevant for your study
with substantial in professionals carin accessible to peop	as developed using participatory design principles, but from patients with T2DM and health g for such patients. Content was designed to be le with a wide range of literacy and health literacy ntial content provided in both video and text."
5-iii) Revisions an	d undating
Revisions and upda application/interver ntervention underw and/or content was	ing. Clearly mention the date and/or version number of the tion (and comparator, if applicable) evaluated, or describe whether the ent major changes during the evaluation process, or whether the development "frozen" during the trial. Describe dynamic components such as news feeds or nich may have an impact on the replicability of the intervention (for unexpected
	1 2 3 4 5
subitem not at all in	portant O O • O essential
Copy and paste rele	ddress subitem 5-iii? vant sections from the manuscript (include quotes in quotation marks "like this" otes from your manuscript), or elaborate on this item by providing additional teems, or briefly explain why the item is not applicable/relevant for your study
nformation not in t	
There has bee no	ance methods on quality assurance methods to ensure accuracy and quality of information

Does your paper add Copy and paste releva	iress subi										
					_		alas I				191
to indicate direct quot information not in the	es from yo	ur m	nanu	ıscr	ipt),	or elabor	ate on thi	is item	by provi	ding addit	ional
information not in the	IIIS, OI DITE	пуе	xpic	1111	wiiy	the item i	з пот арр	nicable	e/Televall	t for your	Study
5-v) Ensure replicab											
screenshots/screen			•			•			_		
Ensure replicability by video, and/or providing											
should in principle be											
	1	2	3	1	5						
	<u>'</u>		3	4	J						
subitem not at all impo	ortant ()	\bigcirc	•	\bigcirc	\bigcirc	essentia	I				
							_				
Copy and paste releva	nt sections	s fro	m th	ne r							
Copy and paste releva to indicate direct quot	ant sections es from yo	s fro ur m	m th	ne r	ipt),	or elabor	ate on thi	is item	by provi	ding addit	ional
Copy and paste releva to indicate direct quot information not in the The website https://w	ant sections les from yo ms, or brie www.help-d	s fro ur m fly e liabe	m th nanu expla etes.	ne n uscr ain v	ipt), why J.uk/	or elabor the item i has been	ate on thi s not app n put into	is item Ilicable	by provi	ding addit	ional
Copy and paste releva to indicate direct quot information not in the The website https://w practice and the gen	ant sections les from yo ms, or brie www.help-d leral inform	s fro ur m fly e liabe ation	m the nanuexpla etes. n ca	ne ruscr ain v org n b	ipt), why J.uk/ e br	or elabor the item i has been owsed wit	ate on thi s not app n put into thout log-	is item Ilicable	by provi	ding addit	ional
Copy and paste releva to indicate direct quot information not in the The website https://w	ant sections les from yo ms, or brie www.help-d leral inform vant to repli	s fro ur m fly e liabe ation cate	m the nanuexplace tes. In calculus the calcu	ne nuscrain v ain v org n b	ript), why g.uk/ e br udy o	or elabor the item i has been owsed wit	ate on thi s not app n put into thout log-	is item Ilicable	by provi	ding addit	ional
Copy and paste relevatory indicate direct quote information not in the The website https://wpractice and the genother researchers wwebsite team and matchis is not a technical	ant sections les from yo ms, or brie www.help-d leral inform vant to repli ake arrang	s fro ur m fly e liabe ation cate eme	m the nanuexplace etes. In calculus the etes. In calculus ethe ent o	ne nuscrain value or one or on	ript), why J.uk/ e br udy o se.	or elabor the item i has been owsed wit could conf	ate on thi s not app n put into thout log- tact the	is item licable in.	by provi	ding addit	ional
Copy and paste relevatory indicate direct quote information not in the The website https://wpractice and the genother researchers website team and material to indicate the control of the	ant sections les from yo ms, or brie www.help-d leral inform vant to repli ake arrang	s fro ur m fly e liabe ation cate eme	m the nanuexplace etes. In calculus the etes. In calculus ethe ent o	ne nuscrain value or one or on	ript), why J.uk/ e br udy o se.	or elabor the item i has been owsed wit could conf	ate on thi s not app n put into thout log- tact the	is item licable in.	by provi	ding addit	ional
Copy and paste relevato indicate direct quotinformation not in the The website https://wpractice and the genother researchers wwebsite team and matchis is not a technical	ant sections les from yo ms, or brie www.help-d leral inform vant to repli ake arrang	s fro ur m fly e liabe ation cate eme	m the nanuexplace etes. In calculus the etes. In calculus ethe ent o	ne nuscrain value or one or on	ript), why J.uk/ e br udy o se.	or elabor the item i has been owsed wit could conf	ate on thi s not app n put into thout log- tact the	is item licable in.	by provi	ding addit	ional
Copy and paste relevatory indicate direct quote information not in the The website https://wpractice and the genother researchers wwebsite team and matchis is not a technical	ant sections les from yo ms, or brie www.help-d leral inform vant to repli ake arrang	s fro ur m fly e liabe ation cate eme	m the nanuexplace etes. In calculus the etes. In calculus ethe ent o	ne nuscrain value or one or on	ript), why J.uk/ e br udy o se.	or elabor the item i has been owsed wit could cont	ate on thi s not app n put into thout log- tact the	is item licable in.	by provi	ding addit	ional
Copy and paste releva to indicate direct quotinformation not in the The website https://w practice and the gen Other researchers website team and mathis is not a technica other publications.	nnt sections res from yo ms, or brie www.help-d reral inform vant to repli ake arrang al article. T	s fro ur m fly e liabe ation cate eme	m the nanuexplace etes. In calculus the etes. In calculus ethe ent o	ne nuscrain value or one or on	ript), why J.uk/ e br udy o se.	or elabor the item i has been owsed wit could cont	ate on thi s not app n put into thout log- tact the	is item licable in.	by provi	ding addit	ional
Copy and paste relevato indicate direct quotinformation not in the The website https://wpractice and the genother researchers wwebsite team and mathis is not a technication other publications.	ant sections les from yo ms, or brie www.help-d leral inform rant to repli ake arrang al article. T	s fro ur m fly e liabe ation cate eme he d	m the captain of the	ne ruscr Jorg Jorg Jorg Jorg Jorg Jorg Jorg Jor	ript), why g.uk/ e br udy (se. n tha	or elabor the item i has been owsed wit could conf at regard i	ate on this not apportunity in put into thout log-tact the s describ	is item ilicable in. e in	by provi e/relevan	ding addit t for your	ional study
practice and the gen- Other researchers we website team and matching is not a technical other publications. 5-vi) Digital preservation: Policy disappear over the content of the process of the content of the preservation of th	ant sections less from yo ms, or brie www.help-d leral inform rant to replicate arrang al article. To ation Provide the learns of the learns of the	s fro ur m fly e liabe ation cate eme he d	m the nanuexplatex of tricks; and the nanuexplatex of tricks	ne ruscrain value in soro	ipt), why j.uk/ e br udy o see. n tha	or elabor the item i has been owsed wit could conf at regard i ication, bu ke sure th	ate on this not apportunity in put into thout log-tact the s describute at as the interver	is item ilicable in. e in	by provi e/relevan	ding addit t for your ikely to ch d (Internei	ional study
Copy and paste releva to indicate direct quotinformation not in the The website https://w practice and the gen Other researchers website team and mathis is not a technica other publications. 5-vi) Digital preservation: Podisappear over the con Archive, webcitation.o	ant sections tes from yoms, or brie www.help-deral inform rant to replicate arrangal article. The artion trovide the larg, and/or part of the arg, and	s fro ur m fly e liabe ation cate eme he d	m th nanu explaintes n ca e the ent o detai	ne ruscr Jorg Jorg Jorg to Jorg to Jor	uk/ g.uk/ g.uk/ e br udy o se. n tha appl mai the s	or elabor the item i has been owsed wit could cont at regard i ication, bu ke sure th source coo	ate on this s not app n put into thout log-tact the s describ ut as the ie interver de or screen	is item ilicable in. e in	ntion is I	ding addit t for your ikely to ch d (Internet s alongsic	ional study ange or de the
Copy and paste relevatory information not in the The website https://w practice and the gen Other researchers website team and mathries is not a technical other publications. 5-vi) Digital preservation: Programmer over the country webcitation. Or article). As pages behite to indicate the country webcitation.	ant sections tes from yoms, or brie mww.help-dueral inform vant to replicate arrangal article. The ation trovide the brg, and/or pind login so	s fro ur m fly e liabe ation cate eme he d	m th nanu explaintes n ca e the ent o detai	ne ruscr Jorg Jorg Jorg to Jorg to Jor	uk/ g.uk/ g.uk/ e br udy o se. n tha appl mai the s	or elabor the item i has been owsed wit could cont at regard i ication, bu ke sure th source coo	ate on this s not app n put into thout log-tact the s describ ut as the ie interver de or screen	is item ilicable in. e in	ntion is I	ding addit t for your ikely to ch d (Internet s alongsic	ional study ange or de the
Copy and paste relevator indicate direct quote formation not in the The website https://w practice and the gen Other researchers website team and mathris is not a technical other publications. 5-vi) Digital preservation: Prodisappear over the contact of the pastern of the contact of the pastern of the pastern of the contact of the pastern of the pa	ant sections tes from yoms, or brie www.help-dueral inform rant to replicate arrangal article. To ation trovide the lurse of the org, and/or pind login so t login.	ur m fly e liabe ation cate eme he d	of t rs; a lishii ns c	ne ruscreament of the control of the	ipt), why j.uk/ e br udy of the see. In the see the see the see. In the see th	or elabor the item i has been owsed wit could cont at regard i ication, bu ke sure th source coo	ate on this s not app n put into thout log-tact the s describ ut as the ie interver de or screen	is item ilicable in. e in	ntion is I	ding addit t for your ikely to ch d (Internet s alongsic	ional study ange or de the
Copy and paste relevatory information not in the The website https://w practice and the gen Other researchers website team and mathries is not a technical other publications. 5-vi) Digital preservation: Programmer over the country webcitation. Or article). As pages behite to indicate the country webcitation.	ant sections tes from yoms, or brie www.help-dueral inform rant to replicate arrangal article. To ation trovide the lurse of the org, and/or pind login so t login.	ur m fly e liabe ation cate eme he d	m th nanu explaintes n ca e the ent o detai	ne ruscreament of the control of the	ipt), why j.uk/ e br udy of the see. In the see the see the see. In the see th	or elabor the item i has been owsed wit could cont at regard i ication, bu ke sure th source coo	ate on this s not app n put into thout log-tact the s describ ut as the ie interver de or screen	is item ilicable in. e in	ntion is I	ding addit t for your ikely to ch d (Internet s alongsic	ional study ange or de the
Copy and paste relevato indicate direct quotinformation not in the The website https://wpractice and the gen. Other researchers wwebsite team and mathis is not a technica other publications. 5-vi) Digital preservation: Page 15-vi	ant sections tes from yoms, or brie mww.help-deral inform to replicate arrangal article. The article of the larg, and/or pind login so togin.	ur m fly e liabe ation cate eme he d	of t of t of t alishin	ne ruscreain vo.organ be stuof usils in the aulso ng teanr	ript), why juk/e brudy of sec. In that the second the s	or elabor the item i has been owsed wit could cont at regard i ication, but ke sure the source coo be archive	ate on this not apportunity in put into thout log-tact the s describut as the interverde or scred, consider	is item ilicable in. e in	ntion is I	ding addit t for your ikely to ch d (Internet s alongsic	ional study ange or de the

"EM is the Managing Director of a not-for-profit Community Interest Company established to disseminate HeLP-Diabetes across the UK (https://www.help-diabetes.org.uk/)."
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).
1 2 3 4 5
subitem not at all important O O essential
information not in the ms, or briefly explain why the item is not applicable/relevant for your study "The intervention consisted of facilitated access to HeLP-Diabetes. Facilitation consisted of an introductory training session with practice nurses. In this appointment patients were shown how to log on, set a user name and password, and introduced to the content of the website." As this study was conducted from the perspective of the NHS and personal and social services in the UK, participants were given free use of the programme. If adopted by the NHS, the access to
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].
1 2 3 4 5
subitem not at all important \bigcirc \bigcirc \bigcirc essential
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

"HeLP-Diabetes was a theoretically informed, web-based programme, whose overall goals were to improve health outcomes and reduce diabetes-related distress. Overall content was guided by the Corbin and Strauss model...Content was designed to be accessible to people with a wide range of literacy and health literacy skills, with all essential content provided in both video and text." Comparator"a simple information website, based on the information readily available in the public domain"

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.

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

"had access to their allocated intervention for 12 months after randomisation"

One of the advantages of the website-based self-management programme is that users can have the flexibility in 'doses'. Although the programme was devided into several genearl topics, it was intended to be used ad libitum.

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).

Does your paper address subitem 5-x?

"The intervention consisted of facilitated access to HeLP-Diabetes. Facilitation consisted of an introductory training session with practice nurses. In this appointment patients were shown how to log on, set a user name and password, and introduced to the content of the website."

"A medical information scientist reviewed the diabetes-related research published each month and provided a summary of important, useful or relevant research."

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).

1 2 3 4 5
subitem not at all important \bigcirc \bigcirc \bigcirc \bigcirc essential

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

"Engagement with the programme was promoted through regular newsletters, emails and mobile text -messages containing updates on latest diabetes-related research or practice, seasonally-relevant advice, and links to specific relevant parts of the programme."

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.

1 2 3 4 5
subitem not at all important \bigcirc \bigcirc \bigcirc \bigcirc essential

Does your paper address subitem 5-xii? *

"The intervention consisted of facilitated access to HeLP-Diabetes. Facilitation consisted of an introductory training session with practice nurses. In this appointment patients were shown how to log on, set a user name and password, and introduced to the content of the website."
6a) Completely defined pre-specified primary and
secondary outcome measures, including how and
when they were assessed
When they were deceded
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 health outcomes of the intervention were assessed through two measurements: diabetes-related distress, measured by the Problem Areas in Diabetes (PAID) questionnaire, andEQ-5D-3Lwere collected online at baseline, three months, and 12 months follow-up."
"The majority of informationwere extracted from participants'
medical records by practice or research nurses. The remainder of the service use data were collectedusing a self-report
6a-i) Online questionnaires: describe if they were validated for online use and apply CHERRIES items to describe how the questionnaires were designed/deployed If outcomes were obtained through online questionnaires, describe if they were validated for online use and apply CHERRIES items to describe how the questionnaires were designed/deployed [9]. 1 2 3 4 5
subitem not at all important O O O essential
Does your paper address subitem 6a-i?
Copy and paste relevant sections from manuscript text

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

ed (lo	ogin	s, lo	gfile	e ana	alysis, etc.)	Use/adopt		netri	cs ar	e imp	ortant	
1	2	3	4	5								
	0	0	•	0	essential							
					•							
how whe	, an n qu	d w lualita	hen ative	qua e fee	alitative fe	edback fro						
					essential							
					int tout							
	sub sub sub	how, an when quorms, interest of the subitent	ed (logins, lould be reported 1 2 3 subitem 6a etions from rency of constant 1 2 3 subitem 6a etions intervier 1 2 3 subitem 6a etions from rency of constant 1 2 subitem 6a etions from rency of constant 1 2 subitem 6a etions from rency of constant 1 2 subitem 6a etions from rency of constant 1 2 subitem 6a etions from rency of constant 1 2 subitem 6a etions from rency of constant 1 2 subitem 6a etions from rency of constant 1 2 subitem 6a etions from rency of constant 1 2 subitem 6a etions from rency of constant 1 2 subitem 6a etions from rency of constant 1 2 subitem 6a etions from rency of constant 1 2 subitem 6a etions from rency of constant 1 2 subitem 6a etions from rency of constant 1 2 subitem 6a etions from rency of constant 1 2 subitem 6a etions from rency o	ed (logins, logificated in the logical subitem 6a-ii? subitem 6a-ii? etions from manuel ency of consultate how, and when when qualitative orms, interviews, 1 2 3 4 subitem 6a-iii?	ed (logins, logfile and uld be reported in an 1 2 3 4 5	ed (logins, logfile analysis, etc.). uld be reported in any ehealth tr 1 2 3 4 5 subitem 6a-ii? etions from manuscript text ency of consultation or episode, when qualitative feedback from orms, interviews, focus groups). 1 2 3 4 5 1 2 3 4 5 2 9 9 9 essential	ed (logins, logfile analysis, etc.). Use/adopuld be reported in any ehealth trial. 1 2 3 4 5 subitem 6a-ii? etions from manuscript text ency of consultation or episode, shown in how, and when qualitative feedback from participar orms, interviews, focus groups). 1 2 3 4 5 1 2 3 4 5 2 3 4 5 3 4 5 3 6 0 0 0 essential	uld be reported in any ehealth trial. 1 2 3 4 5 subitem 6a-ii? etions from manuscript text ency of consultation or episode, shown in when qualitative feedback from participants was orms, interviews, focus groups). 1 2 3 4 5 1 2 3 4 5	ed (logins, logfile analysis, etc.). Use/adoption metriculd be reported in any ehealth trial. 1 2 3 4 5 subitem 6a-ii? ctions from manuscript text ency of consultation or episode, shown in when qualitative feedback from particular partic	ed (logins, logfile analysis, etc.). Use/adoption metrics are uld be reported in any ehealth trial. 1 2 3 4 5 subitem 6a-ii? etions from manuscript text ency of consultation or episode, shown in when qualitative feedback from participant when qualitative feedback from participants was obtained bruss, interviews, focus groups). 1 2 3 4 5 1 2 3 4 5 2 9 0 0 essential	ed (logins, logfile analysis, etc.). Use/adoption metrics are impuld be reported in any ehealth trial. 1 2 3 4 5 subitem 6a-ii? ctions from manuscript text ency of consultation or episode, shown in how, and when qualitative feedback from participants was when qualitative feedback from participants was obtained (e.gorms, interviews, focus groups). 1 2 3 4 5 1 2 3 4 5 2 0 0 0 essential	ed (logins, logfile analysis, etc.). Use/adoption metrics are important uld be reported in any ehealth trial. 1 2 3 4 5 subitem 6a-ii? etions from manuscript text ency of consultation or episode, shown in how, and when qualitative feedback from participants was when qualitative feedback from participants was obtained (e.g., orms, interviews, focus groups). 1 2 3 4 5 1 2 3 4 5 2 0 0 0 essential

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

Does your paper address CONSORT subitem 6b? *

"There was no changes to the methods after the protocol was agreed and the start of the trial"	

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.

	1	2	3	4	5	
subitem not at all important	\bigcirc	\bigcirc	\bigcirc	•	\bigcirc	essentia

Does your paper address subitem 7a-i?

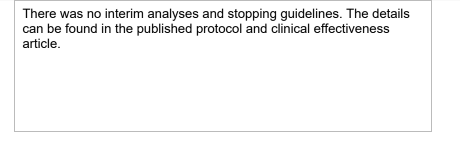
Copy and paste relevant sections from manuscript title (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

The economic outcomes were not the primary outcome of the trial, therefore the sample size was not calculated based on costs or QALYs.

The details of the calculation of the sample size could be found in the published protocol and clinical effectiveness article. The calculation for the primary outcome took into account expected 85% follow-up rate at 12 months.

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

Does your paper address CONSORT subitem 7b? *



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

"individually randomised to either intervention or control group using a web-based randomisation independently of the trial team."

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

"Randomisation was conducted in a 1:1 ratio using random permuted blocks of sizes 2, 4 and 6, stratified by recruitment centre."

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

"Eligible participants were briefed on the trial by practice nurse or our study team (Patient Information Sheet see Multimedia Appendix 2) and then individually randomised to either intervention or control group using a web-based randomisation independently of the trial team. Randomisation was conducted in a 1:1 ratio using random permuted blocks of sizes 2, 4 and 6, stratified by recruitment centre."

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

"Eligible participants were briefed on the trial by practice nurse or our study team (Patient Information Sheet see Multimedia Appendix 2) and then individually randomised to either intervention or control group using a web-based randomisation independently of the trial team. Randomisation was conducted in a 1:1 ratio using random permuted blocks of sizes 2, 4 and 6, stratified by recruitment centre."

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).

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

Does your paper address subitem 11a-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 were informed the trial compared two forms of webbased support but were blinded as which was the intervention and which the comparator."

Nurses provided the facilitation couldn't be blinded. The analyst of this analysis didn't have access to data before the trial ended. On conducting the analysis, it was not possible to blind the analyst as the cost of intervention had to be allocated to the intervention group.

11a-ii) Discuss e.g., whether participants knew which intervention was the "intervention of interest" and which one was the "comparator"

Informed consent procedures (4a-ii) can create biases and certain expectations - discuss e.g., whether participants knew which intervention was the "intervention of interest" and which one was the "comparator".

	1	2	3	4	5	
subitem not at all important	\bigcirc	•	\bigcirc	\bigcirc	\bigcirc	essential

Does your paper address subitem 11a-ii?

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

We provided an information website as comparator so that the participants wouldn't suspect which was the "intervention of interest".

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? *

They were both websites.	

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

"A linear mixed effects model was fitted with the 12-month outcome as the dependent variable, adjusting for the baseline variables age, sex, presence of pre-existing cardiovascular disease, duration of diabetes and smoking status and corresponding baseline outcome (costs over 12 months before baseline, PAID score at baseline and EQ-5D-3L utility value at baseline, respectively) as fixed effect terms. Centre effects were included as random-effects in the analysis. No time dependent

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

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

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

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

"Multiple imputation was used as the primary method to account for missing data in both baseline and follow-up. A chained equation model was developed and predictive mean matching was used as the imputation method for continuous variables, using the five nearest neighbours to the prediction as a set to draw from. All missing data were imputed separately by trial group. The imputation was performed on the aggregated level estimate"

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

"A complete case analysis was undertaken to assess the performance of the imputation model compared to a complete case analysis that assumes missing completely at random. The intervention cost estimated in the primary analysis was based on trial participants only, which would be unrealistic when implementing to a wider population with T2DM. We therefore undertook a one-way sensitivity analysis, exploring the cost per user as numbers of users increase if rolled out in the community."

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

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

X26-i) Comment on ethics committee approval

Does your paper address subitem X26-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

"Ethics approval was obtained from Camden and Islington National Research Ethics Service (NRES) committee, reference 12/LO/1571."

x26-ii) Outline informed consent procedures

Outline informed consent procedures e.g., if consent was obtained offline or online (how? Checkbox, etc.?), and what information was provided (see 4a-ii). See [6] for some items to be included in informed consent documents.

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

Does your paper address subitem X26-II?	
Copy and paste relevant sections from the manuscript (include quotes in to indicate direct quotes from your manuscript), or elaborate on this item information not in the ms, or briefly explain why the item is not applicable	by providing additional
The informed consent procedure has been described in the protocol	
and the clinical effectiveness article.	
X26-iii) Safety and security procedures	
Safety and security procedures, incl. privacy considerations, and any step	
likelihood or detection of harm (e.g., education and training, availability o	f a hotline)
1 2 3 4 5	
subitem not at all important \bigcirc \bigcirc \bigcirc \bigcirc essential	
Does your paper address subitem X26-iii?	
Copy and paste relevant sections from the manuscript (include quotes in to indicate direct quotes from your manuscript), or elaborate on this item	
information not in the ms, or briefly explain why the item is not applicable	
This was a self-management website programme for people with	
diabetes. It was unlikely to cause any safety issue.	

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? *

"Of 374 participants randomised, 185 were allocated to the intervention and 189 to the control group"
13b) For each group, losses and exclusions after randomisation, together with reasons
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 "The analysis followed a pre-specified analysis plan, comparing the groups as randomised (intention-to-treat)." There was no exclusions after randomisation.
The losses in this case were the ones who had missing values on the variables taken into account in the analysis. "the complete case analysis was performed on 197 participants (96 in the intervention group, 101 in the control group)."
13b-i) Attrition diagram Strongly recommended: An attrition diagram (e.g., proportion of participants still logging in or using the intervention/comparator in each group plotted over time, similar to a survival curve) or other figures or tables demonstrating usage/dose/engagement.
1 2 3 4 5
subitem not at all important \bigcirc \bigcirc \bigcirc \bigcirc 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 This was presented elsewhere. As explained above, given a large amount of variables required, attrition diagram was not practical nor meaningful.

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

"Recruitment took place between September 2013 and December 2014."

There was no set date for follow-up only relative to the individual randomisation date.

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"

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

There was no noticeable such events fell into the study period. However, even if there were, it should only be discussed in the discussion or as a reason to change the methods in the methods section.

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

Does your paper address CONSORT subitem 14b? *

,	showing baseline demographic and
clinical cha	racteristics for each group
	able, a description of care providers (case volume, qualification, expertis volume) in each group
Does your paper a	ddress CONSORT subitem 15? *
to indicate direct que	vant sections from the manuscript (include quotes in quotation marks "like th otes from your manuscript), or elaborate on this item by providing additional ne ms, or briefly explain why the item is not applicable/relevant for your study
This was presented	d in the clinical effectiveness article, which the manuscript. It would increase the length of the
manuscript even fu	
15-i) Report demo	graphics associated with digital divide issues
n ehealth trials it is	particularly important to report demographics associated with digital divide
ssues, sucn as age, of the participants, if	education, gender, social-economic status, computer/Internet/ehealth literac f known.
	1 2 3 4 5
subitom not at all im	nportant O O O essential
subitem not at an im	- essential
	ddress subitem 15-i? *
	vant sections from the manuscript (include quotes in quotation marks "like th otes from your manuscript), or elaborate on this item by providing additional
	ne ms, or briefly explain why the item is not applicable/relevant for your study
nformation not in th	singled to everying the demonstration consists of
nformation not in the The trial wasn't des with digital divide is	signed to examine the demographics associated ssues. Nor was it secondary outcome. Some
The trial wasn't des with digital divide is statistics could be	

16) For each group, number of participants (denominator) included in each analysis and whether the analysis was by original assigned groups

16-i) Report multiple "denominators" and provide definitions

Report multiple "denominators" and provide definitions: Report N's (and effect sizes) "across a range of study participation [and use] thresholds" [1], e.g., N exposed, N consented, N used more than x times, N used more than y weeks, N participants "used" the intervention/comparator at specific pre-defined time points of interest (in absolute and relative numbers per group). Always clearly define "use" of the intervention.

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

Does your paper address subitem 16-i? *

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

"The analysis followed a pre-specified analysis plan, comparing the groups as randomised (intention-to-treat)."

This analysis didn't report number of participants 'used' the intervention because the economic evaluation is intended to reflect the reality, especially the costs. In practice, not everyone will comply with the treatment regime to the same extent. The running costs of the website wouldn't change with number of use.

16-ii) Primary analysis should be intent-to-treat

Primary analysis should be intent-to-treat, secondary analyses could include comparing only "users", with the appropriate caveats that this is no longer a randomized sample (see 18-i).

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

Does your paper address subitem 16-ii?

"The analysis followed a pre-specified analysis plan, comparing the
groups as randomised (intention-to-treat)."

17a) For each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval)

Does	your	paper	address	CONSORT	subitem	17a? *
-------------	------	-------	---------	----------------	---------	--------

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 standard diviations, standard errors, and 95% confidence intervals were presented in the relevant tables.	

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

In addition to primary/secondary (clinical) outcomes, the presentation of process outcomes such as metrics of use and intensity of use (dose, exposure) and their operational definitions is critical. This does not only refer to metrics of attrition (13-b) (often a binary variable), but also to more continuous exposure metrics such as "average session length". These must be accompanied by a technical description how a metric like a "session" is defined (e.g., timeout after idle time) [1] (report under item 6a).

	1	2	3	4	5	
subitem not at all important	\bigcirc	\bigcirc	•	\bigcirc	\bigcirc	essential

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

As mentioned before, one of the advantage of the web-based self-management programme was that it didn't have an intended dosage.

17b) For binary outcomes, presentation of both absolute and relative effect sizes is recommended

Does your paper address CONSORT subitem 17b? *

to indicate direct quotes	sections from the manuscript (include quotes in quotation marks "like this" from your manuscript), or elaborate on this item by providing additional s, or briefly explain why the item is not applicable/relevant for your study
There was no binary ou	tcomes.
19) Populte of	f any other analyses performed
*	f any other analyses performed,
	group analyses and adjusted analyses, pre-specified from exploratory
Does your paper addre	ss CONSORT subitem 18? *
to indicate direct quotes	sections from the manuscript (include quotes in quotation marks "like this" from your manuscript), or elaborate on this item by providing additional s, or briefly explain why the item is not applicable/relevant for your study
reported. There were pr	and a one-way sensitivity analysis were re-specified analyses.
A subgroup analysis of co	s of comparing only users comparing only users is not uncommon in ehealth trials, but if done, it must self-selected sample and no longer an unbiased sample from a
(111	1 2 3 4 5
subitem not at all importa	ant O O • O essential
Does your paper addre	ss subitem 18-i?
to indicate direct quotes	sections from the manuscript (include quotes in quotation marks "like this" from your manuscript), or elaborate on this item by providing additional or briefly explain why the item is not applicable/relevant for your study
	om an economic point of view, comparing he purpose of the study.

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 in the ms, or briefly explain why the item is not applicable/relevant for your study
This is a web-based self-management programme so there was no harms or unintended effects expected or detected.
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 \bigcirc \bigcirc \bigcirc 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 There was no such breaches occurred.
19-ii) Include qualitative feedback from participants or observations from

staff/researchers

Include qualitative feedback from participants or observations from staff/researchers, if available, on strengths and shortcomings of the application, especially if they point to unintended/unexpected effects or uses. This includes (if available) reasons for why people did or did not use the application as intended by the developers.

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

Does your paper address subitem 19-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
As a quantitative analysis, qualitative feedback was not part of this manuscript.
manuscript.
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
and unequal expertise of care providers of centers in each group
22-i) Restate study questions and summarize the answers suggested by the data, starting
with primary outcomes and process outcomes (use) Restate study questions and summarize the answers suggested by the data, starting with primary
outcomes and process outcomes (use).
1 2 3 4 5
subitem not at all important O O essential
Does your paper address subitem 22-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
"In this within-trial economic evaluation, we found that a web-based
self-management intervention, HeLP-Diabetes, was highly likely to be cost-effective when compared to the WTP threshold
recommended by NICE, even with a small number of users. Once there were over 363 users, HeLP-Diabetes became dominant (i.e.
less costly and more effective) compared to a free information-only
website such as that provided by NHS Choices or Diabetes UK."

22-ii) Highlight unanswered new questions, suggest future research

Highlight unanswered new questions, suggest future research.

1 2 3 4 5

subitem not at all important	\bigcirc	\bigcirc	\bigcirc	ledow	\bigcirc	essential
------------------------------	------------	------------	------------	-------	------------	-----------

Does your paper address subitem 22-ii?

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

"As with any research, one of the main questions is if the results of the study can be generalised to a wider population."..."Further research is needed on how digital health interventions such as HeLP-Diabetes can be developed, delivered and maintained in a sustainable and cost-effective manner."..."Currently, the successful realisation of this effect might lie in identifying the more susceptible user groups and engaging them first. More empirical studies would be needed establish the pathway from research to practice."

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

20-i) Typical limitations in ehealth trials

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

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

Does your paper address subitem 20-i? *

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

"only the current medication at the time of data collection was extracted and the assumption was made that all medications were for chronic use." "...not to include the investment costs incurred in the development of the intervention in our analysis...there may be more maintenance costs over a longer period of time as technology changes continually, and there may be a need for software updates...not to undertake long-term modelling, based on the observed improvement in HbA1c"

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

study results for other organ	Γset	ting	and			ar, discuss I patient p								
	1	2	3	4	5									
subitem not at all important	: 0	0	0	•	0	essential								
Does your paper address Copy and paste relevant sec to indicate direct quotes fro information not in the ms, o "We do not expect interne	ction m your brie t-bas	s fro our r efly o	om t nan expl nter	he nuscrain vent	ipt), why tions	or elabora the item is to be suit	te on this not applicable for	iten cabl	n k	by p	prov	iding	g addit	ional
everyone, but with the inte is potential for digital healt on health care systems in "Currently, the successful identifying the more susce first."	th int the l reali	erve ong satio	ntio run on o	ns to ." f this	o he s eff	lp alleviate ect might l	the burde e in							
21-ii) Discuss if there we application setting Discuss if there were eleme (e.g., prompts/reminders, mand what impact the omissi intervention is applied outsi	nts i nore l	n the	e RC an i	CT th nvol	nat w lvem	ould be di	fferent in a	a ro	uti r o	ine	app er co	olicat o-inte	tion se erventi	etting ons)
		_	_											
	1	2	3	4	5									
subitem not at all important						essential								

OTHER INFORMATION

23) Registration number and name of trial registry

to indicate direct quotes from your manuscript), or elaborate on this item information not in the ms, or briefly explain why the item is not applicable	
ISRCTN02123133	

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

24) Where the full trial protocol can be accessed, if available

Does your paper address CONSORT subitem 24? *

Does your paper address CONSORT subitem 23? *

Cite a Multimedia Appendix, other reference, or 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

doi: 10.1186/s12913-015-1246-9 One of the references in the manuscript.	

25) Sources of funding and other support (such as supply of drugs), role of funders

Does your paper address CONSORT subitem 25? *

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

inis project was funded by the National Institute for Health
Research (NIHR)"
"additionally funded by the UK Medical Research Council
(MR/L003120/1), British Heart Foundation (RG/13/13/30194) and
UK National Institute for Health Research Cambridge Biomedical
Research Centre "

"The funders had no role in the study design, data collection, data analysis, data interpretation, or writing of the report and the

X27) Conflicts of Interest (not a CONSORT item)

X27-i) State the relation addition to the usual study team towards the from or identical with	ne system	bein	ig ev	/alu	ated	, i.e., state i	f the autho	, .	
	1	2	3	4	5				
subitem not at all imp	ortant O	0	0	•	0	essential			
D		••	- V0	· - :	0				
Does your paper add Copy and paste releva to indicate direct quot Information not in the	ant section tes from yo	s fro	om t nani	he r uscr	nanı ipt),	or elaborat	e on this it	em by provi	ding additional
"AF is an NIHR Seni Oxford Biomedical F				d red	ceive	es funding f	rom NIHR		
EM is the Managing				or-p	rofit	Community	Interest		
Company establishe (https://www.help-dia	ed to disse	mina	ate F						
(Intps://www.neip-die	abeles.org	j.uk/	,.						
About the C	ONSC	R	TE	ΞΗ	IE/	ALTH (check	ist	
As a result of using	this chec								pt? *
As a result of using yes, major change	this chec								pt? *
As a result of using yes, major change yes, minor change	this chec								pt? *
As a result of using yes, major change yes, minor change no	this chec s	klist	t, di	d yo	ou m	nake chanç	jes in you	r manuscri	
As a result of using yes, major change yes, minor change no	this chec s s	klist	t, die	d yo	ou m	nake chanç	jes in you	r manuscri	
As a result of using yes, major change yes, minor change no What were the most	this chec s s	klist	t, die	d yo	ou m	nake chanç	jes in you	r manuscri	
As a result of using yes, major change yes, minor change no What were the most	this chec s s	klist	t, die	d yo	ou m	nake chanç	jes in you	r manuscri	
As a result of using yes, major change yes, minor change no What were the most	this chec s s	klist	t, die	d yo	ou m	nake chanç	jes in you	r manuscri	
As a result of using yes, major change yes, minor change no What were the most	this chec s s	klist	t, die	d yo	ou m	nake chanç	jes in you	r manuscri	
As a result of using yes, major change yes, minor change no What were the most	this chec s s	klist	t, die	d yo	ou m	nake chanç	jes in you	r manuscri	
As a result of using yes, major change yes, minor change no What were the most Added more details	this checes s importar in the met	klist	aang	d yo	you	made as a	es in you	r manuscri	checklist?
About the C As a result of using yes, major change yes, minor change no What were the most Added more details How much time did in your manuscript	this checes s importar in the met	klist	aang	d yo	you	made as a	es in you	r manuscri	checklist?
As a result of using yes, major change yes, minor change no What were the most Added more details How much time did in your manuscript	this checes s importar in the met	klist	aang	d yo	you	made as a	es in you	r manuscri	checklist?
As a result of using yes, major change yes, minor change no What were the most Added more details How much time did in your manuscript	this checes s importar in the met	klist	aang	d yo	you	made as a	es in you	r manuscri	checklist?
As a result of using yes, major change yes, minor change no What were the most Added more details How much time did in your manuscript	this checes s importar in the met	klist	aang	d yo	you	made as a	es in you	r manuscri	checklist?

yes	
O no	
Other:	
Would you like to become involved in the	CONSORT FHEALTH group?
	olved in participating in a workshop and writing an
○ yes	
ono no	
Other:	
Any other comments or questions on CO	NSORT EHEALTH
probably is the methods for cost-effectivened quite some room. If all details of RCT must	
with this checklist, the manuscript would be journal article.	•
with this checklist, the manuscript would be journal article. STOP - Save this form as To generate a record that you filled in this	•
with this checklist, the manuscript would be journal article. STOP - Save this form as To generate a record that you filled in this page (on a Mac, simply select "print" and the same that you fill the same	PDF before you click submit
with this checklist, the manuscript would be journal article. STOP - Save this form as To generate a record that you filled in this page (on a Mac, simply select "print" and to when you submit your (revised) paper to stile. Don't worry if some text in the textboxes in the sextboxes in the sextboxes in the sextboxes.	PDF before you click submit form, we recommend to generate a PDF of this then select "print as PDF") before you submit it.
with this checklist, the manuscript would be journal article. STOP - Save this form as To generate a record that you filled in this page (on a Mac, simply select "print" and the work of the substitution of the same of the	PDF before you click submit form, we recommend to generate a PDF of this hen select "print as PDF") before you submit it. JMIR, please upload the PDF as supplementary
with this checklist, the manuscript would be journal article. STOP - Save this form as To generate a record that you filled in this page (on a Mac, simply select "print" and the When you submit your (revised) paper to viile. Don't worry if some text in the textboxes in our database. Thank you! Final step: Click submit!	PDF before you click submit form, we recommend to generate a PDF of this hen select "print as PDF") before you submit it. JMIR, please upload the PDF as supplementary secut off, as we still have the complete information
with this checklist, the manuscript would be journal article. STOP - Save this form as To generate a record that you filled in this page (on a Mac, simply select "print" and the When you submit your (revised) paper to viile. Don't worry if some text in the textboxes in our database. Thank you! Final step: Click submit!	PDF before you click submit form, we recommend to generate a PDF of this hen select "print as PDF") before you submit it. JMIR, please upload the PDF as supplementary secut off, as we still have the complete information
with this checklist, the manuscript would be journal article. STOP - Save this form as To generate a record that you filled in this page (on a Mac, simply select "print" and the when you submit your (revised) paper to stille. Don't worry if some text in the textboxes in our database. Thank you! Final step: Click submit! Click submit so we have your answers in the state of the submit is the submit in the submit is the submit in the submit is t	S PDF before you click submit form, we recommend to generate a PDF of this hen select "print as PDF") before you submit it. JMIR, please upload the PDF as supplementary is cut off, as we still have the complete information our database!

Report Abuse - Terms of Service - Additional Terms