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

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

The goal of the CONSORT EHEALTH checklist and guideline is to be a) a guide for reporting for authors of RCTs, b) to form a basis for appraisal of an ehealth trial (in terms of validity)

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

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

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

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

Mandatory reporting items are marked with a red *. In the textboxes, either copy & paste the relevant sections from your manuscript into this form please include any quotes from your manuscript in QUOTATION MARKS, or answer directly by providing additional information not in the manuscript, or elaborating on why the item was not relevant for this study.

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

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

Citation Suggestion (if you append the pdf as Appendix we suggest to cite this paper in the caption):
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.imir.org/2011/4/e126/
doi: 10.2196/jmir.1923
PMID: 22209829

* Required

Your name *

Jeffrey Lambert

Primary Affiliation (short), City, Country * University of Toronto, Toronto, Canada

University of Exeter, Exeter, UK

Your e-mail address * abc@gmail.com

j.d.lambert@exeter.ac.uk

Title of your manuscript *

Provide the (draft) title of your manuscript. A pilot randomized controlled trial of an administratively supported web-based intervention using behavioral activation and physical activity for adults with depression: The eMotion study. Name of your App/Software/Intervention * If there is a short and a long/alternate name, write the short name first and add the long name in brackets. eMotion **Evaluated Version (if any)** e.g. "V1", "Release 2017-03-01", "Version 2.0.27913" Language(s) * What language is the intervention/app in? If multiple languages are available, separate by comma (e.g. "English, French") **English** URL of your Intervention Website or App * e.g. a direct link to the mobile app on app in appstore (itunes, Google Play), or URL of the website. If the intervention is a DVD or hardware, you can also link to an Amazon page. Your answer URL of an image/screenshot (optional) Your answer Accessibility * Can an enduser access the intervention presently? access is free and open access only for special usergroups, not open access is open to everyone, but requires payment/subscription/in-app purchases app/intervention no longer accessible Other: Primary Medical Indication/Disease/Condition * e.g. "Stress", "Diabetes", or define the target group in brackets after the condition, e.g. "Autism (Parents of children with)", "Alzheimers (Informal Caregivers of)" Depression Primary Outcomes measured in trial * comma-separated list of primary outcomes reported in the trial Depression (PHQ-8)

噩

Secondary/other outcomes

Are there any other outcomes the intervention is expected to affect?

Objective physical activity, self reported physical activity, Anxiety
Recommended "Dose" * What do the instructions for users say on how often the app should be used?
Approximately Daily
Approximately Weekly
Approximately Monthly
Approximately Yearly
O "as needed"
Other:
Approx. Percentage of Users (starters) still using the app as recommended after 3 months *
unknown / not evaluated
0-10%
O 11-20%
O 21-30%
O 31-40%
O 41-50%
O 51-60%
O 61-70%
71%-80%
81-90%
91-100%
Other:
Overall, was the app/intervention effective? *
yes: all primary outcomes were significantly better in intervention group vs control
 partly: SOME primary outcomes were significantly better in intervention group vs control
O no statistically significant difference between control and intervention
O potentially harmful: control was significantly better than intervention in one or more outcomes
inconclusive: more research is needed
Other: This is a pilot study. Exploractory analysis found an effect in favour

Article Preparation Status/Stage * At which stage in your article preparation are you currently (at the time you fill in this form)
onot submitted yet - in early draft status
not submitted yet - in late draft status, just before submission
Submitted to a journal but not reviewed yet
O submitted to a journal and after receiving initial reviewer comments
Submitted to a journal and accepted, but not published yet
O published
Other:
Journal * If you already know where you will submit this paper (or if it is already submitted), please provide the journal name (if it is not JMIR, provide the journal name under "other")
O not submitted yet / unclear where I will submit this
Journal of Medical Internet Research (JMIR)
JMIR mHealth and UHealth
○ JMIR Serious Games
○ JMiR Mental Health
○ JMIR Public Health
JMiR Formative Research
Other JMIR sister journal
Other:
Is this a full powered effectiveness trial or a pilot/feasibility trial?
Pilot/feasibility
O Fully powered
Manuscript tracking number * If this is a JMIR submission, please provide the manuscript tracking number under "other" (The ms tracking number can be found in the submission acknowledgement email, or when you login as author in JMIR. If the paper is already published in JMIR, then the ms tracking number is the four- digit number at the end of the DOI, to be found at the bottom of each published article in JMIR)
no ms number (yet) / not (yet) submitted to / published in JMIR
Other:
TITLE AND ABSTRACT
1a) TITLE: Identification as a randomized trial in the title
1a) Does your paper address CONSORT item 1a? * Le does the title contain the phrase "Randomized Controlled Trial"? (if not, explain the reason under "other")
yes
Other:

廽

1a-i) Identify the mode of delivery in the title Identify the mode of delivery. Preferably use "web-based" and/or "mobile" and/or "electronic game in the title. Avoid ambiguous terms like "online", "virtual", "interactive". Use "Internet-based" only if Intervention includes non-web-based internet components (e.g. email), use "computer-based" or "electronic" only if offline products are used. Use "virtual" only in the context of "virtual reality" (3-D worlds). Use "online" only in the context of "online support groups". Complement or substitute product names with broader terms for the class of products (such as "mobile" or "smart phone" instead of "iphone"), especially if the application runs on different platforms. subitem not at • essential all important Does your paper address subitem 1a-i? * Copy and paste relevant sections from manuscript title (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for "web-based" 1a-ii) Non-web-based components or important co-interventions Mention non-web-based components or important co-interventions in title, if any (e.g., "with telephone support"). subitem not at 0 0 0 0 essential all important Does your paper address subitem 1a-ii? Copy and paste relevant sections from manuscript title (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study N/A 1a-iii) Primary condition or target group in the title Mention primary condition or target group in the title, if any (e.g., "for children with Type I Diabetes") Example: A Web-based and Mobile Intervention with Telephone Support for Children with Type I Diabetes: Randomized Controlled Trial subitem not at 0 essential all important Does your paper address subitem 1a-iii? * Copy and paste relevant sections from manuscript title (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study "for adults with depression" 1b) ABSTRACT: Structured summary of trial design,

NPT extension: Description of experimental treatment, comparator, care providers, centers, and blinding status.

methods, results, and conclusions

1b-i) Key features/functionalities/components of the										
intervention and comparator in the METHODS section of the										
ABSTRACT										
Mention key features/ abstract. If possible, a the needs of systemat report in the abstract v body of text, consider	lso mention ic reviewer what the ma	theories and and indexe	d principles	used for deling imports	esigning the int synonym:	site. Keep in mind s. (Note: Only				
			196							
subitem not at ail important	0	0	0	0	•	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										
eMotion is an administratively supported weekly modular programme aiming to help people use key behavior change techniques (e.g., graded tasks, action planning and self-monitoring) to re-engage in routine, pleasurable or necessary activities, with a focus on physical activities. Feasibility data was collected which included: recruitment and trial retention rates, the fidelity of intervention delivery, receipt and enactment, and the acceptability of the intervention and data collection procedures."										
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)										
subitem not at all important	0	0	•	0	0	essential				
Does your par Copy and paste releve "like this" to indicate additional information your study	ant section	s from the m	anuscript a	bstract (inc t), or elabor	ate on this it	em by providing				
"administratively	supporte	d"								

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)

0	0	•	0	0	essentia
	0	0 0	0 0 0	0 0 0 0	0 0 0 0

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

"Data was collected for the primary (depression) and secondary outcomes (e.g. anxiety, PA, fidelity and client satisfaction) at baseline and two months post randomization using self-reported online questionnaires and accelerometers."

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)

Does your paper address subitem 1b-iv?

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

Within the intervention group, the median number of logins, modules accessed and total minutes spent on eMotion was 3 (IQR = 2.0 to 8.0), 3 (IQR = 2.0 to 5.0) and 41.3 (IQR = 18.9 to 90.4) respectively.

1b-v) CONCLUSIONS/DISCUSSION in abstract for negative trials

Conclusions/Discussions in abstract for negative trials: Discuss the primary outcome - if the trial is negative (primary outcome not changed), and the intervention was not used, discuss whether negative results are attributable to lack of uptake and discuss reasons. (Note: Only report in the abstract what the main paper is reporting. If this Information is missing from the main body of text, consider adding it)

subitem not at all important	0	0	③	0	0	essential

Does your paper address subitem 1b-v?

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

"Exploratory data showed that PHQ-8 levels were reduced more for the intervention group than the control group at two months post randomization (Adjusted Mean Difference: -3.6 (95% CI: -6.1 to -1.1)."

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)

subitem not at all important	0	0	0	0	•	essential

Does your paper address subitem 2a-i? *

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

*Depression has a significant detrimental impact on individuals and their families, as well as increased utilisation of health services, and reduced productivity at work leading to considerable economic costs" "Physical activity (PA) has also been shown to be effective in treating (Cooney et al., 2013) and preventing (Mammen & Faulkner, 2013) depression, and is often cited by patients as their preferred treatment option (Searle et al., 2011; Ussher, Stanbury, Cheeseman, & Faulkner, 2007)." "Web-based interventions could provide a useful way of overcoming these limitations by delivering such interventions outside of existing services, recruiting directly from the community and standardising fidelity (Lambert, Greaves, Farrand, Cross, et al., 2017; Watkins et al., 2016)."

2a-ii) Scientific background, rationale: What is known about the

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.

subitem not at all important	0	0	(a)	0	0	essentia
all lillpurtant						

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

"There is a paucity of studies that have attempted to deliver web-based interventions promoting PA for depression (Carlbring et al., 2013; Nyström et al., 2017; Rosenbaum, Newby, Steel, Andrews, & Ward, 2015; Soucy, Provencher, Fortler, & McFadden, 2017), and no studies have explored the feasibly of delivering a web based intervention combining BA and PA for people with depression. "

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 purpose of the present study is to examine the feasibility of delivering an online intervention (eMotion) combining PA with BA to people for depression, and to explore its effects on depression and PA."

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 eMotion trial was a two-arm, individually randomized, parallel group pilot RCT with a nested process evaluation"

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

Does your paper address CONSORT subitem 3b? *

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

"The eMotion trial provided 'minimal contact' administrative support (Farrand & Woodford, 2013) at week two of the intervention to provide the participant with a rationale for the use of self-help materials and check-ins related to progress (but with no focus on any clinical or behavior change issues). This support was initially intended to be provided by an independent 'supporter'. However due to resource issues, this support was provided by the lead author."

3b-i) Bug fixes, Downtimes, Content Changes

Bug fixes, Downtimes, Content Changes: ehealth systems are often dynamic systems. A description of changes to methods therefore also includes important changes made on the intervention or comparator during the trial (e.g., major bug fixes or changes in the functionality or content) (5-lii) and other "unexpected events" that may have influenced study design such as staff changes, system failures/downtimes, etc. [2].

all important	0	0	•		0	essential
Does your particle Copy and paste relevation indicate direct quo information not in the	ant sections tes from yo	s from the ur manus	manuscript cript), or elab	(include quot orate on this	item by pro	viding additional
N/A						
1476						
4a) Eliaibility	anitania :	£		. 17		
4a) Eligibility	criteria	ror par	ucipant	5 /		
Does your par	oer addr	ess CO	NSORT	subitem	427 *	
Copy and paste relev						ion marks "like th
to indicate direct quo	tes from yo	ur manusc	cript), or elab	orate on this	item by pro	viding additional
information not in the	ms, or brie	rıy explain	why the iten	n is not applic	cable/releva	nt for your study
Yes						
4a-i) Compute						
Computer / Internet is explicitly clarified.	iteracy is oft	ten an imp	licit "de fact	o" eligibility c	riterion - this	should be
explicitly clarified.						
subitem not at	\sim	\sim		_	~	
all important	\cup	\circ	•	\circ	\circ	essential

Does your paper address subitem 4a-i?

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

"were computer literate"

4a-ii) Open vs. Open vs. closed, web- (online vs. offline), e.g. web-based trial, or the assessment), i.e., to clarify if participants whether technical or to detect/prevent the	based vs. fa g., from an o ere were fac what degree were quasi-a logistical me	nce-to-face as pen access ve-to-face cor got the study anonymous a	ssessments website or fr nponents (a y team to kn and whether	: Mention h om a clinic s part of th ow the part having mul	ow participar , and clarify if e interventior ticipant. In on tiple identitie	nts were recruited this was a purely or for line-only trials, s was possible or
subitem not at all important	0	0	0	0	•	essential
Does your pap Copy and paste releva to Indicate direct quo information not in the	ant sections tes from you ms, or brief	from the ma ir manuscrip ily explain wh	nuscript (in t), or elabora ny the item is	clude quote ate on this s not applic	item by provi able/relevan	ding additional t for your study
"Eligible adults we newspapers, soci banners on webs	ial media (e.g. Facet	ook supp	ort group	s, Twitter)	ents in weekly and through
4a-iii) Informal Information given duthe informed consense also item X26), a and may also blas re-	ring recruitm t procedures s this inform	ent. Specify (e.g., publis	how particip	pants were led consent	t documentat	ion as appendix,
subitem not at all important	0	0	0	0	•	essentiai
Does your pai Copy and paste relev to indicate direct quo information not in the	ant sections	from the manuscrip	anuscript (in ot), or elabor	ciude quot rate on this	item by provi	ding additional
"After contacting sent the participa online screening they could withdo obliged to provid complete the onliby the lead author author contacted	ant inform questionr raw from the a reason line conse or to assess the partic	ation shee naire, via e the study a n. Once pa nt form fo as particip cipant via	et (PIS), comail. At the any time of the any t	onsent for his point, to e without read the a screen lity. After clarify the	rm, and a li they were i consequer PIS, they w ing question screening, e study pro-	nk to the informed that ince or being were asked to innaire used the lead cedures and
The participant v return it in a pre- to wear an accel measures were t	vas instru stamped a erometer	cted to we and addres did not pre	ar for the ssed enve eclude ran	accelero lope. Hov domizati	meter for s vever, parti on. Further	even days and cipant refusal baseline
Participants were						_
4b) Settings						ected
Does your pa Copy and paste rele to indicate direct qu information not in the	vant section	s from the m	nanuscript (i	nclude quo	tes in quotati item by prov	iding additional
Ves						

pased trials) or othe	rwise.	(seit-)asse	essea throug	n online que:	stionnaires (as common in wei
subitem not at all important	0	0	0	0	•	essential
Does your pa copy and paste relet o indicate direct qualiformation not in th	vant section otes from ye e ms, or bri	ns from the our manusc efly explain	manuscript cript), or elab why the iter	(include quo orate on this n is not appli	item by pro cable/relev	viding additional ant for your study
delivered at scre ompleted version				-randomiz	ation usin	g an online self
1b-ii) Report I deport how institution ffiliations with presi with regards to an in	nal affiliatio	ons are dispoitals or uni	played to por iversities ma	ential partici y affect volu	pants (on el nteer rates,	nealth media], as use, and reactions
subitem not at all important	0	0	•	0	0	essential
OOCS YOUR PAP opy and paste releve indicate direct quo	ant sections	s from the	manuscript (include quot	es in quotat	ion marks "like this
formation not in the	ms, or brie	fly explain	why the iten	is not appli	cable/releva	nt for your study
I/A						
/A) The interve llow replicati	ntions t	for eacl uding h	h group now and	with suf when th	ficient c	letails to actually
) The interve llow replicati dministered	on, incl	uding h	now and	when th	ey were	actually
The interve llow replicati dministered -i) Mention n ponsors, and ention names, cred ention serve serve	ames, compension of the compen	uding heredenties	ial, affilia e developers of the softw	when the ations of ations, sponsors, a lare, this nee	the dev	e actually velopers,
) The interve llow replicati dministered -i) Mention n ponsors, and ention names, cred ention names, cred ention names or cred interest" section or	ames, compension of the compen	uding heredenties	ial, affilia e developers of the softw	when the ations of ations, sponsors, a lare, this nee	the dev	e actually velopers,
The interve llow replicati dministered i) Mention n ponsors, and ention names, cred athors/evaluators are interest" section or subitem not at all important Oes your pap	ames, commens of owners of mentioned	ess sub	ial, affilial e developers of the softwin the manu	ations of ations of ations of ations of ations of ations of ations ation	the development of the developme	relopers, 6) (if lared in a "Conflic" essential
) The interve llow replicati dministered -i) Mention n ponsors, and ention names, cred thors/evaluators ar interest" section or ubitem not at all important oes your pap	ames, commens of the	ess sub from the nur manuscrify explain was develored.	ial, affilial e developers of the softwin the manuscript (i ipt), or elabowhy the item	etions of ations of ations of a sponsors, a sare, this need script).	the development of the developme	elopers, felopers, f
The interve llow replication of the property o	ames, commens, commens, commens, commens, commens, commens, commentioned commentioned commentioned commens, com	ess sub from the nur manuscrify explain via as developments. LTTF) or	ial, affilial edevelopers of the softwin the manuscript (ilipt), or elabowhy the Item oped by the nline platfor elopmer of the applica	etions of ations of ations of a sponsors, a sare, this need script). ? nclude quote rate on this is not applice estudy autorm was usent processition and prevention and p	the devand owners in quotatitem by provable/relevanthors (***, seed to hossious formatics of the service of the	elopers, folif essential essential on marks "like this iding additional at for your study **, **, ** & **). st the

Does your paper address subitem 5-ii?

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

"eMotion was adapted from the BAcPAc intervention (Farrand et al., 2014; Pentecost et al., 2015), using the Centre for eHealth Research and Disease Management (CeHReS) roadmap (van Gemert-Pijnen et al., 2011). The CeHReS roadmap is intended to help the planning, coordination and execution of the participatory development process of eHealth. In eMotion, this involved using patient public involvement, usability testing and a structured literature search. A full description of the eMotion intervention and its developmental process is provided elsewhere (Lambert, Greaves, Farrand, Haase, & Taylor, 2017)."

5-iii) Revisions and updating

Revisions and updatin application/interventic intervention underwen and/or content was "fi changing content white events see item 3b).	on (and com at major cha rozen" durir	nparator, if a inges during ng the trial. I	applicable) (g the evalua Describe dy	evaluated, or tion process namic comp	describe who controls who describe who describes who describes with the describes and describes who describes who describes with the describes who describes and describes who describes and describes who describes and describes who describes and describes	the development as news feeds o
subitem not at all important	0	0	•	0	0	essential
Does your pap Copy and paste releve to indicate direct quot information not in the	ant sections tes from vo	from the muscri	nanuscript (ipt), or elabo	nclude quot orate on this	item by prov	iding additional
N/A						
		-	*			
5-iv) Quality as Provide information of provided [1], if applica	n quality as	e meth	ods ethods to er	nsure accura	cy and quali	ty of information
subitem not at all important	0	0	•	0	0	essential
Does your par Copy and paste relev to indicate direct quo information not in the	ant sections	s from the r	nanuscript (ript), or elab	include quo orate on this	item by pro	viding additional
N/A						
5-v) Ensure re	nlicahil	ity by n	ublishin	a the so	urce co	de. and/or
providing scre	enshol	s/scree	en-captu	re video	, and/or	providing
flowcharts of Ensure replicability by video, and/or providi should in principle be	the algorithms the publishing flowchar	orithms the source	used code, and/ gorithms us	or providing	screenshots	s/screen-capture er researchers
subitem not at all important	0	0	•	0	0	essential
Does your pa Copy and paste relev to indicate direct que	vant section	s from the	manuscript	(include qua	otes in quota s item by pro	tion marks "like t

information not in the ms, or briefly explain why the item is not applicable/relevant for your study

Yes: Found in intervention description and published paper on interventiomn development (Lambert et al., 2017)

5-vi) Digital pro			ennlication	hut ae the in	tervention !	s likely to change o
disappear over the co Archive, <u>webcitation.o</u> article). As pages beh are accessible withou	urse of the rg, and/or ind login s	years; also publishing t	make sure the source o	the intervent code or scree	ion is archiv enshots/vide	red (Internet eos alongside the
subitem not at all important	0	0	•	0	0	essential
Does your pap Copy and paste releva to indicate direct quot information not in the	int section es from yo	s from the r	nanuscript (ript), or elab	(include quo orate on this	item by pro	viding additional
Screenshots prov	ided In L	ambert et	al., (2017	"		
5-vii) Access Access: Describe how pay (or were paid) or r						
how participants obta editors/reviewers/rea reviewers/readers to o	ined "acce ders, cons	ss to the pla ider to provi	atform and l de a "backd	nternet" [1]. loor" login a	To ensure a count or de	ccess for mo mode for
subitem not at all important	0	0	•	0	0	essential
Does your pap Copy and paste releva to indicate direct quot information not in the	int section es from yo	s from the r our manuscr	nanuscript (ript), or elab	include quo orate on this	item by pro	viding additional
"Through a series 1), eMotion teach						•
activities that proving reinforcement. eM						
modules, 1 gener!		•				
consisting of visu opens. Printable, !					-	
to the slides to all	ow dowr	nloading t	o a perso	nal compu	iter or and	ther device
(e.g. tablet or sma week by the eMot						sent once a
5-viii) Mode of		-				
the interventio						
comparator, and the to behaviour change tect	hniques, po	ersuasive fe	atures, etc.,	see e.g., [7,	8] for termin	ology). This
includes an in-depth of developed it) [1]," whe track their progress at delivery channels and	ther (and h	now] it is tail feedback" [lored to indi 6]. This also	vidual circur includes a c	nstances an description o	d allows users to of communication

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

0

0

0

essential

0

hyperlinks to other resources, etc. [1].

subitem not at

all important

Does your paper address subitem 5-viii? *

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

"The eMotion intervention is a web-based course that provides people with access to an evidence-based treatment based on BA with added PA promotion. People with depression commonly reduce activities which they perceive as burdensome, making less effort to do things they may have previously enjoyed. By reducing such activities, people with depression experience temporary relief which then negatively reinforces the likelihood of avoiding further activities. However, avoiding such activities has a long term cost, as it reduces the opportunity for positive reinforcement which occurs when people engage in social and personal activities which bring them pleasure and achievement. PA is an example of such an activity which people with depression often avoid but

has the potential t health benefits. "	o provide	e additior	ial anti-de	epressive be	enefits as	well as added
5-ix) Describe Describe use paramet instructions or recommendations of use, if an	ers (e.g., in mendations	ntended "do s were give	ses" and op n to the use	r, e.g., regardi		
subitem not at all important	0	0	•	0	0 7	essential
Does your pap Copy and paste releva to indicate direct quot information not in the	nt sections es from yo	s from the r ur manusci	nanuscript ipt), or elab	(include quote orate on this i	tem by pro	iding additional
"Key content relati	ing to the	e rational	e of BA is	front loade	d in the i	ntroduction,
week 1 and week		-	-			
and designed to s						
people to review to		-	•		-	
to log in (e.g. by u		•				,
5-x) Clarify the Clarify the level of hun assistance) in the e-in involved, if any, as wel it is initiated, and the r distinguish between the involvement required in generalizability).	nan involve tervention il as "type o nedium by ne level of l	ement (care or as co-int of assistance which the a human invo	providers of ervention (of se offered, to assistance in livernent rec	or health profe detall number he timing and is delivered", h quired for the t	and experti frequency t may be ne trial, and the	se of professional of the support, how cessary to e level of human
	(8)	50				
subitem not at all important	0	0	•	0	0	essential
Does your pap Copy and paste releva to indicate direct quot information not in the	nt section: es from yo	s from the r our manusci	manuscript ript), or elab	(include quote orate on this	tem by pro	viding additional
"brief administrati provided at week		ational s	upport vla	a a 10 minu	te phone	call was
					8	
5-xi) Report an Report any prompts/rv SMS) to use the applic between the level of p for a routine application	eminders u cation, wha rompts/rei	ised: Clarify at triggered minders rec	if there we them, frequ juired for th	re prompts (le ency etc. It m e trial, and the	ay be neces level of pr	ssery to distinguisl compts/reminders
subitem not at all important	0	0	0	0	0	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

"Automated reminder emails are also sent once a week by the eMotion programme following registration."

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.

Does your paper address subitem 5-xii? *

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

N/A

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

Does your paper address CONSORT subitem 6a? *

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

Yes

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].

subitem not at all important O O O essential

Does your paper address subitem 6a-i?

Copy and paste relevant sections from manuscript text

"The PHQ-8 has been shown to have good validity, reliability, sensitivity and specificity (Kroenke et al., 2009) and been used in previous web-based intervention studies of low mood and depression (Baumeister et al., 2014; Richards & Richardson, 2012)"

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

Describe whether and how "use" (including intensity of use/dosage) was defined/measured/monitored (logins, logfile analysis, etc.). Use/adoption metrics are important process outcomes that should be reported in any ehealth trial.

subitem not at all important O O O o essential

Does your paper address subitem 6a-ii?

Copy and paste relevant sections from manuscript text

"Delivery fidelity was assessed using website usage statistics from the online intervention database. This database provided individual level data about whether the participant registered for eMotion, modules accessed and the total time spent on each module. Participant receipt and enactment of the intended intervention processes were measured using online questionnaires. For fidelity of receipt, two conceptual approaches were used, The first focused on participants' understanding of how emotions, behaviors, thoughts and physical feelings affect each other to maintain depression over time. A single item was used based on questions used in a previous study (Brawley, Arbour-Nicitopoulos, & Martin Ginis, 2013). The item employed a five-point Likert response scale (Strongly Agree to Strongly Disagree) assessing participant agreement with the following statement: "I understand how emotions, behaviors, thoughts and physical feelings affect each other to maintain depression over time." A second measure of receipt assessed participants' perceived ability to use the intended behavior change techniques (BCTs). This was assessed by asking participants to rate their confidence in using specific BCTs (i.e. identification of suitable activities, grading activities for ease of use, planning and dealing with setbacks) in the last two months on a scale from 1 (not at all confident) to 10 (very confident). This measure was adapted from measures of confidence used in the ProActive trial (Hardeman et al., 2008). Finally, to assess enactment, we asked participants if they had used specific BCTs relating to behavioral activation (BA) In the last two months using a binary scale (yes/no). This measure was adapted from similar measures of BCT usage that showed that enactment was significantly associated with weight loss, providing initial evidence of the validity of this type of measure (Hankonen et al., 2014). '

6a-iii) Describe whether, how, and when qualitative feedback from participants was obtained

Describe whether, how, and when qualitative feedback from participants was obtained (e.g., through emails, feedback forms, interviews, focus groups).

subitem not at all important	0	0	0	\odot	essentia

Does your paper address subitem 6a-iii? Copy and paste relevant sections from manuscript text

"The Client Satisfaction Questionnaire – Short Form (CSQ-SF) is a 4 item measure to assess participant satisfaction regarding their use of eMotion two months post-randomization (given to intervention participants only). This measure was administered using an online questionnaire and has been used to assess treatment satisfaction in web-based studies of other interventions for depression (Williams et al., 2013)."

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

Does your paper address CONSORT subitem 6b? *
Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study

N/A

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

subitem not at all important O O O essential

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

"Due to the pilot nature of the study, no formal sample size calculations were conducted. However, to ensure a suitably reliable estimate of the standard deviations to power a future trial with 90% power, at least 15 people per arm are recommended if the expected effect size lies between 0.3 and 0.7 (Whitehead, Julious, Cooper, & Campbell, 2015). A previous meta-analysis of computer-based psychological treatments for depression reported a moderate effect size (0.56) and a drop-out rate of 57% (Richards & Richardson, 2012). As such, a target sample size of 62 (accounting for a possible attrition rate of 50%) to ensure at least 15 people per arm at follow up was adopted."

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

ind stopping guidelines

Does your paper address CONSORT subitem 7b? *

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

N/A

18

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

"Once participants completed the baseline assessment, they were randomly allocated to either the intervention or control group using simple randomization at the individual level in a 1:1 ratio and an online randomization service (Sealed Envelope Ltd. 2016). Personal details were anonymized through the use of participant numbers which were entered into the website by the lead author in a consecutive manner (In the order of completed baseline assessment), and the randomization service allocated them to either group A (eMotion) or group B (waiting list)."

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

"Once participants completed the baseline assessment, they were randomly allocated to either the Intervention or control group using simple randomization at the individual level in a 1:1 ratio"

9) Mechanism sequence (sudescribing an interventions	ch as se y steps t	quentia taken to	ily numb	ered co	ntainers	s),
Does your paper Copy and paste relevito indicate direct quo information not in the	ant sections tes from you	from the m	anuscript (in pt), or elabor	clude quote ate on this it	s in quotation em by provi	ding additional
Your answer						
10) Who gene enrolled parti interventions	erated th cipants,	e rando and wh	om alloca o assigr	ation se led parti	quence, icipants	who to
Does your pal Copy and paste relev to indicate direct que information not in the	ant sections	from the m	nanuscript (ir ipt), or elabor	clude quote ate on this i	s in quotation tem by prov	iding additional
"online randomiz	ation serv	ice (Seale	ed Envelop	e Ltd. 201	6)"	
11a) If done, interventions those assess	(for exa	mple, p omes)	articipa and how	nts, care	provid	
NPT: Whether or no	ot administe	ering co-int	erventions v	vere blinde	d to group a	assignment
11a-i) Specify Specify who was blit participants [1, 3] (the assessors, those do	nded, and wh	no wasn't. U clearly ack	Isually, in wel nowledged).	o-based trial but it may b	s it is not po e possible to	o blind outcome
subitem not at all important	0	0	0	0	•	essential
Does your pa Copy and paste rele to indicate direct qu information not in th	vant section	s from the o	manuscript (i ript), or elabo	nclude quot rate on this	item by pro	viding additional
"Due to limited r condition each p nature of the int allocation. How surveys, there w participant's res introduce subje	participant ervention, ever, as ou vas a reduc sponses, o	was allo it was als itcome m ced chanc r for the l	cated follo so impossi leasures w ce of the le ead author	wing rand ble to blin ere taken ad author to misint	lomization d particip using onli influencia erpret res	n. Due to the ants to group ine self-report ng the ponses or

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.,

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".

subitem not at all important	0	0	0	0	•	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

"Due to limited resources for the study, the lead author was not blinded to which condition each participant was allocated following randomization. Due to the nature of the intervention, it was also impossible to blind participants to group allocation. However, as outcome measures were taken using online self-report surveys, there was a reduced chance of the lead author influencing the participant's responses, or for the lead author to misinterpret responses or introduce subjective bias into recorded observations (Edwards, 2010)."

11b) If relevant, description of the similarity of interventions

(this item is usually not relevant for ehealth trials as it refers to similarity of a placebo or sham intervention to a active medication/intervention)

Does your paper address CONSORT subitem 11b? *

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

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

"Quantitative methods were used to explore: the recruitment and attrition rates of trial participants, the feasibility of data collection and study procedures, baseline data (including levels of PA and baseline differences between groups and between dropouts), and fidelity of delivery. Descriptive statistics were produced for all outcomes by trial arm at baseline and two-month follow-up. All quantitative analyses were conducted using Stata SE statistical software release 14 (StataCorp. 2015; College Station, TX). No formal hypothesis-testing relating to primary outcomes was planned as this was a pilot study. However, descriptive statistics were used to assess recruitment and retention rates, and baseline PA levels. Baseline demographic and clinical characteristics were presented descriptively as proportions or as means with standard deviations. Two types of exploratory analysis of the primary outcome for the main trial (PHQ-8) were conducted: 1) linear regression models to report changes in depression with 95% confidence Intervals around the between-group mean difference and 2) logistic regression models which dichotomised the primary outcome to reflect clinically meaningful change (a reduction to below ten on the PHQ-8 indicates that the person may no longer qualify for major depression (Kroenke et al., 2009))."

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]).

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

"The analyses were conducted on an intention-to-treat basis with complete data only, which included those who began treatment and provided follow-up data regardless of treatment compliance. Missing data were not imputed with data from participants who provided data at both time points analysed."

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

"Similar analyses were conducted for anxiety, and objective and self-reported PA. We conducted sensitivity analyses linear regression models to examine the effects of receiving psychological therapies, as well as any substantial differences in baseline characteristics on the findings."

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



X26-i) Comment on ethics committee approval

cubitom not at	_	_	_	\sim	•	
subitem not at all important	O	\circ	\circ	\circ		essentia
an miportant						

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

"This study was approved by the University of Exeter Sports and Health Sciences Research Ethics Committee (AM160316-21 151021/B/03)."

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.

subitem not at	\circ	\circ	\circ	\circ	\circ	essentia
subitem not at all important	\circ	0		0		

Does your paper address subitem X26-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

"Once participants read the PIS, they were asked to complete the online consent form (indicating consent using a checkbox) followed by a screening questionnaire used by the lead author to assess participant eligibility."

X26-iii) Safety and security procedures

Safety and security procedures, incl. privacy considerations, and any steps taken to reduce the likelihood or detection of harm (e.g., education and training, availability of a hotline)

subitem not at all important O O O o essential

Does your paper address subitem X26-iii?

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

"One possible ethlcal issue in this study was suicide risk in people experiencing depression. As this was a research study on a non-clinical sample, all participants were advised on the PIS that the study was not a clinical or NHS treatment and that the University and researchers could not take clinical responsibility for the treatment of any conditions they might have including depression. They were also signposted to other appropriate resources in case they wished to seek formal treatment. If, at any point in the study (e.g. whilst on the phone to a researcher during screening or after inclusion), participants indicated suicidal intent, the University of Exeter Mood Disorders Suicide Risk Protocol was invoked."

RESULTS

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

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

Does your paper address CONSORT subitem 13a? *

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

"A total of 183 people responded to the adverts, with 100 completing screening for eligibility (Figure 1). Of the 183 individuals who initially inquired about the study 100 were still interested screened for eligibility and 62 (34% of those initially enquiring (95% CI: 27 to 41%) and 62% of those who were screened (95% CI: 52 to 71%)) were eligible for inclusion and randomized into the trial between May 2016 and February 2017 (32 eMotion, 30 control)."

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

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

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

*Overall attrition (in relation to the planned main trial primary outcome (PHQ-8) at two months post-randomization was 12 people (19%; 95% CI: 11 to 31%). Of those randomized, 58 (94%) participants provided complete secondary outcome baseline measurements (e.g. GAD7, IPAQ-SF) and 52 (84%) provided usable accelerometer data at baseline. At two-month follow-up, 50 (81% of those randomized (95% CI: 71 to 91 %) provided PHQ-8 (and other survey data) and 24 (39% of those randomized (95% CI: 27 to 52%) provided valid accelerometer data (with >10 hours data for at least 3 weekdays and one weekend day). Mean weekend days of wear time for the accelerometers was 2.0 (SD = 0.2) at baseline and 1.9 (SD = 0.3) at two-month post-randomization. Mean weekday days of wear time for the accelerometers was 4.8 (SD = 0.3) at baseline and 4.6 (SD = 0.9) at two months post randomization. Only 47 (76%) and 33 (53%) participants provided valid IPAQ-SF data at baseline and two months postrandomization respectively. This lack of usable IPAQ-SF data was due to people providing invalid responses to the IPAQ-SF."

13b-i) Attrition Strongly recommende the intervention/comp figures or tables demo	ed: An attriti parator in e	ion diagram ach group p	lotted over	time, similar	icipants still to a survival	logging in or using curve) or other
subitem not at all important	0	0	•	0	0	essential
Does your pap Copy and paste releva (include quotes in que elaborate on this item item is not applicable	ant sections otation mar n by providi	s from the n ks "like this ng additions	nanuscript o " to indicate al informatio	or cite the flg direct quote	s from your	manuscript), or
Your answer						
14a) Dates de	efining t	the perio	ods of r	ecruitme	ent and	follow-up
Does your pap Copy and paste relev to indicate direct quo information not in the	ant section	s from the r	manuscript ript), or elab	(include quot orate on this	es in quotat item by pro	viding additional
Reported in 'fidel	lty' sectio	n				1.4
14a-i) Indicate period						
resources available	or "changes	is reli into i	er hardware	or Internet d	elivery resou	ırces"
subitem not at all !mportant	0	0	•	0	0	essentia!
Does your pa Copy and paste relet to indicate direct quinformation not in the	vant section	ns from the	manuscript	(include quo borate on thi	s item by pro	oviding additional
N/A						

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

Copy and paste relev to indicate direct quo information not in the	ant section tes from yo	s from the our manusc	manuscript ript), or elab	orate on this	tes in quotat item by pro	viding additional
N/A						
15) A table sh characteristic				graphic	and clin	ical
NPT: When applicat etc.) and centers (ve				ers (case vo	lume, qualit	ication, expertise,
Does your par Copy and paste relev- to indicate direct quo information not in the	ant section tes from yo	s from the our manusc	manuscript ript), or elab	(include quo orate on this	tes in quotat item by pro	viding additional
Table 2						
15-i) Report do In ehealth trials it is p issues, such as age, e of the participants, if	articularly i education, g	mportant to	report dem	ographics a	ssociated w	th digital divide
subitem not at all important	0	0	•	0	0	essential
Does your pap Copy and paste releva to indicate direct quoi information not in the	ant sections tes from yo	s from the r ur manusc	manuscript (ript), or elab	include quot orate on this	item by pro	viding additional
Table 2						
16) For each g included in ea original assigi	ch anal	ysis an				
16-i) Report m Report multiple "deno range of study particij than x times, N used r specific pre-defined ti clearly define "use" of	minators" a pation [and more than y me points o	and provide use] thresh weeks, N p of interest (definitions: nolds" [1], e.ç participants	Report N's (g., N exposed "used" the in	and effect si d, N consent tervention/c	zes) "across a ed, N used more omparator at
subitem not at all important	0	0	•	0	0	essential
Does your pap Copy and paste releva to indicate direct quot information not in the	nt sections es from yo	from the nur manuscr	nanuscript (ipt), or elab	include quot orate on this	item by prov	iding additional
Table 3/4		1.00				•
16-ii) Primary a Primary analysis shou "users", with the appro	ld be intent	-to-treat, se	econdary an	alyses could	include corr	
subitem not at	0	0	•	0	0	essential
an important						

Does your paper address subitem 16-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

Analysis were Intend to treat with complete data only

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

"Exploratory analyses (table 4) showed that at two months post randomization, the intervention group had a larger reduction in depressive symptoms than the control group (Adjusted Mean D!ff -3.6, 95% CI: -6.1 to -1.1)."

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).

subitem not at	\circ	\circ	\circ	\circ		essentia
all important	\circ	0	O	O	•	essentia

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

"In total, 28 (90%) of the 32 intervention participants registered for eMotion and clicked on the introduction module. The median number of logins, modules accessed and total minutes spent on eMotion was 3 (IQR = 2.0 to 8.0), 3 (IQR = 2.0 to 5.0) and 41.3 (IQR = 18.9 to 90.4) respectively. 61% of participants completed at least the introduction, week 1 and week 2 and 29% of participants completed up to at least week 4. Only one participant used every module. At two months post-randomization, participants randomized to the eMotion group reported a significant difference, compared to the control group, in levels of understanding about how thoughts, feelings and behaviors affect mood (Adjusted Mean Diff 0.5, 95% Cl: -0.0 to -1.0). Significant differences were also found for confidence to identify (Adjusted Mean DIff 1.4, 95% CI: 0.0 to 2.8), select (Mean Diff 1.3, 95% CI: -0.02 to 2.6) and plan (Mean Diff 1.8, 95% CI: 0.5 to 3.1) achievable activities to improve mood as well as confidence to deal with setbacks (Mean Diff 1.5, 95% CI: 0.2 to 2.7). Those who were randomized to the eMotion group and who endorsed 'no' on the enactment questionnaires at baseline were significantly more likely to select (N = 25; OR: 10, 95% CI: 1.6 to 62.7) and plan (N = 33; OR: 10.3, 95% CI: 2.0 to 52.6) activities to improve their mood at two month follow up.

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



Does your paper address CONSORT subitem 17b? *

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

"In the intervention group, 56% (14/25) of depression scores went below the threshold of 10 on the PHQ-8, compared to 28% (7/25) in the control group (OR 3.3, 95% CI: 1.0 to 10.6)."

18) Results of any other analyses performed, including subgroup analyses and adjusted analyses, distinguishing pre-specified from exploratory



Does your paper address CONSORT subitem 18? *

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

"When receipt of other psychological therapies was entered into the regression analysis as a covariate, the impact of co-treatment was not significant, and the intervention group still had a higher reduction in depressive symptoms than the control group (Mean Diff -3.3, 95% CI: -5.9 to -0.7). Other baseline covariates (age, gender, employment, education level, anti-depressant usage) that may have influenced depression scores were also entered in the regression model together. Findings indicated that none of these variables had a significant covariate effect on depression scores and the residual difference between groups was still significant (Mean Diff -3.1, 95% CI: -5.7 to -0.5). Within the intervention group, linear regression analyses revealed no significant relationships between numbers of modules accessed, number of logins or total minutes spent on the website with depression outcomes. The pattern of change scores within each group is shown in figure 2."

18-i) Subgroup analysis of comparing only users

A subgroup analysis of comparing only users is not uncommon in ehealth trials, but if done, it must be stressed that this is a self-selected sample and no longer an unbiased sample from a randomized trial (see 16-iii).

. 1.24						
subitem not at	\circ	\sim	0	_	_	
all important	\circ	\circ	(9)	\circ	\cdot	essential

Does your paper address subitem 18-i?

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

19) All important harms or unintended effects in each group



(for specific guidance see CONSORT for harms)

Does your paper address CONSORT subitem 19? *

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

No harms or unintended effects observed



19-i) Include p Include privacy breach participants, but also i and other unexpected effects [2].	nes, technica Incidents su	d problems	s. This does elved or real	not only incl privacy brea	ude physical ches [1], tec	hnical problems,
subitem not at all important	0	0	•	0	0	essential
Does your pap Copy and paste releva to indicate direct quoi information not in the	ant sections	from the n	nanuscript (ript), or elab	include quot orate on this	item by prov	iding additional
No privacy breach	nes, techn	ical prob	lems obs	erved		
19-ii) Include observations Include qualitative fer on strengths and sho unintended/unexpect did not use the applications.	from sta edback from intcomings o ted effects o	off/rese participar f the appli r uses. Thi	earchers nts or obser- cation, espe is includes (rations from cially if they f available) r	staff/resear	chers, if available,
subitem not at all important	0	0	•	0	0	essential
Does your pal Copy and paste relev to indicate direct quo information not in the	rant sections otes from yo e ms, or brie	from the ur manusc fly explain	manuscript cript), or elab why the iter	(include quo orate on this n is not appl	item by pro icable/releva	viding additional
Qualitative data	gathered, l	out w!ll b	e reporte	d elsewher	e	
DISCUSSION	7					
22) Interpreta and harms, a	ation co nd cons	nsister idering	nt with re	esults, b elevant e	alancing evidence	benefits
NPT: In addition, ta and unequal expen	ike into acc tise of care	ount the o	choice of th or centers	e comparat in each gro	or, lack of o up	r partial blinding,
22-i) Restate suggested by process outc Restate study quest outcomes and proce	the dat omes (L	a, start ise) mmarize th	ting with	primary	outcon	nes and
subitem not at all important	0	0	0	0	•	essential
Does your pa Copy and paste rele to indicate direct qu information not in the	evant section	s from the	e manuscrip script), or ela	(include que borate on th	is item by pr	ation marks "like this" oviding additional vant for your study
"The present stu eMotion interve elevated depres concerning the	ntion. We slve symp	success toms. T	fully recru he trial als	ited a less to had acc	active po	pulation with

22-ii) Highlight	t unansv	wered ne	ew ques	tions, su	iggest fi	uture
research	4-					
Highlight unanswered	new questi	ons, suggest	tuture resea	arch.		
subitem not at all important	0	0	0	0	•	essential
Does your pap Copy and paste releva to indicate direct quot information not in the	nt sections es from you	from the ma r manuscript	nuscript (inc t), or elabora	clude quotes ite on this ite	em by provid	ling additional
Reported in discus	ssion "Imp	olications 1	for future	research"	section	
20) Trial limita imprecision, a						bias,
20-i) Typical III Typical limitations in e often look at a multipli non-use of the interver unexpected events.	health trials	s: Participant omes, increa	s in ehealth sing risk for	trials are rar a Type I erre	or. Discuss l	piases due to
subitem not at all important	0	0	0	0	•	essential
Does your pap Copy and paste releva to indicate direct quot information not in the	nt sections es from you	from the mai r manuscript	nuscript (inc t), or elabora	lude quotes te on this ite	m by provid	ing additional
Reported in discus	ssion "Stre	engths and	limitation	ns" section	1	3
21) Generalisa trial findings	bility (e	xternal v	/alidity,	applicat	oility) of	the
NPT: External validity patients, and care pro		•	-		ntion, comp	arators,
21-i) Generaliz. Generalizability to othe population, outside of study results for other	er population a RCT settir	ns: In particu ng, and gener	lar, discuss	generalizabi		
subitem not at ali important	0	0	•	0	0	essential
Does your pape Copy and paste releval to indicate direct quote information not in the	nt sections t es from you	from the mar manuscript	nuscript (inc), or elabora	lude quotes te on this ite	m by provid	ing additional
Reported in discus	ssion "Rel	ationship 1	to other lit	erature" s	ection	
21-ii) Discuss i different in a ro Discuss if there were e (e.g., prompts/reminde and what impact the or intervention is applied	Outine a elements in t ers, more hu mission of t	pplication The RCT that The man involved The se elemen	on settin would be dif ment, trainin ts could hav	g iferent in a re g sessions o	outine applic or other co-i	cation setting nterventions)
subitem not at all important	0	0	•	0	0	essential

Does your	paper	address	subitem	21-ii?
-----------	-------	---------	---------	--------

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

Reported in discussion "Strengths and limitations" section

OTHER INFORMATION

23) Registration number and name of trial registry

Does your paper address CONSORT subitem 23? *

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

ClinicalTrials.gov NC-T03084055

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

Does your paper address CONSORT subitem 24? *

Cite a Multimedia Appendix, other reference, or copy and paste relevant sections from the 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

N/A

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

In acknowledgements section

X27) Conflicts of Interest (not a CONSORT item)

X27-i) State the relation of the study team towards the system being evaluated

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

subitem not at 0 0 0 0 essential all important

Does your paper address subitem X27-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 additiona information not in the ms, or briefly explain why the item is not applicable/relevant for your study

No conflicts of interest

BB

About the CONSORT EHEALTH checklist

12/02/2018

manuscript? *
O yes, major changes
yes, minor changes
O no
What were the most important changes you made as a result of using this checklist?
Adding further detail around participant flow
How much time did you spend on going through the checklist INCLUDING making changes in your manuscript *
Approx 2 hours
As a result of using this checklist, do you think your manuscript has improved? $\mbox{\ensuremath{^{\star}}}$
yes
O no
Other:
Would you like to become involved in the CONSORT EHEALTH group? This would involve for example becoming involved in participating in a workshop and writing an "Explanation and Elaboration" document
(a) yes
O no
Other:
Any other comments or questions on CONSORT EHEALTH
Your answer
STOP - Save this form as PDF before you click submit
To generate a record that you filled in this form, we recommend to generate a PDF of this page (on a Mac, simply select "print" and then select "print as PDF") before you submit it.
When you submit your (revised) paper to JMIR, please upload the PDF as supplementary file.
Don't worry if some text in the textboxes is cut off, as we still have the complete information in our database. Thank you!
Final step: Click submit!
335.333.333.33
Click submit so we have your answers in our database!
SUBMIT
Never submit passwords through Google Forms

This content is neither created not endorsed by Google. Report Abuse - Terms of Service - Additional Terms

Google Forms