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

PMID: 22209829
* Required
Your name *
First Last
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edward-rojas@uiowa.edu
edward-rojas@drowa.edd
Title of your manuscript *
Provide the (draft) title of your manuscript.
Acceptance and Commitment Therapy Delivered via a Mobile Phone Messaging Robot
Decreases Postoperative Opioid Utilization in Orthopaedic Trauma Patients: Randomized Control Trial
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.
ACT-Based Mobile Phone Messages

doi: 10.2196/jmir.1923

Evaluated Version (if any)
e.g. "V1", "Release 2017-03-01", "Version 2.0.27913"
Your answer
Language(s) *  What language is the intervention/app in? If multiple languages are available, separate by comma (e.g. "English, French")  English
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
Tour 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: Access to the intervention software is granted through a login portal signup cro

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)"
postoperative pain
Primary Outcomes measured in trial * comma-separated list of primary outcomes reported in the trial
opioid tablet utilization
Secondary/other outcomes  Are there any other outcomes the intervention is expected to affect?
patient reported outcome measures
Recommended "Dose" * What do the instructions for users say on how often the app should be used?
Approximately Daily
Approximately Weekly
Approximately Monthly
Approximately Yearly
as needed"
Other: Twice-daily mobile phone messages were delivered to patients

Approx. Percentage of Users (starters) still using the app as recommended after 3 months *
unknown / not evaluated
0-10%
O 11-20%
21-30%
31-40%
O 41-50%
51-60%
61-70%
71%-80%
81-90%
91-100%
Other: intervention is only for 2 weeks following surgery.
Overall, was the app/intervention effective? *
yes: all primary outcomes were significantly better in intervention group vs control
partly: SOME primary outcomes were significantly better in intervention group vs control
on statistically significant difference between control and intervention
outcomes potentially harmful: control was significantly better than intervention in one or more
inconclusive: more research is needed
Other:

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

Is this a full powered effectiveness trial or a pilot/feasibility trial? *  Pilot/feasibility
Fully powered
Manuscript tracking number *  If this is a JMIR submission, please provide the manuscript tracking number under "other" (The ms tracking number can be found in the submission acknowledgement email, or when you login as author in JMIR. If the paper is already published in JMIR, then the ms tracking number is the four-digit number at the end of the DOI, to be found at the bottom of each published article in JMIR)
no ms number (yet) / not (yet) submitted to / published in JMIR  Other: 17750
TITLE AND ABSTRACT
1a) TITLE: Identification as a randomized trial in the title
<ul><li>1a) Does your paper address CONSORT item 1a? *</li><li>I.e does the title contain the phrase "Randomized Controlled Trial"? (if not, explain the reason under "other")</li></ul>
<ul><li>yes</li><li>Other:</li></ul>

1a-i) Identify the mode of de Identify the mode of delivery. Prefera title. Avoid ambiguous terms like "onlincludes non-web-based Internet comoffline products are used. Use "virtua only in the context of "online support terms for the class of products (such application runs on different platform"	bly use "wine", "virtunponents (l' only in groups".	reb-based' ual", "inter (e.g. email the contex Compleme	and/or "n active". Us ), use "cor t of "virtua ent or subs	e "Interne nputer-ba al reality" ( stitute pro	t-based" or sed" or "ele (3-D worlds duct name	nly if Intervention ectronic" only if e). Use "online" s with broader
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1a-ii) Non-web-based components support").	or import	ant co-int	erventions	in title, if	any (e.g., "	
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Does your paper address sub Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e	m manuso uscript), o	cript title (i or elaborat	e on this i	tem by pro	viding add	itional

1a-iii) Primary condition or target group in the title  Mention primary condition or target group in the title, if any (e.g., "for children with Type I Diabetes")  Example: A Web-based and Mobile Intervention with Telephone Support for Children with Type I Diabetes:  Randomized Controlled Trial								
	1	2	3	4	5			
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Does your paper address su	hitem 1:	a-iii? <b>*</b>						
Copy and paste relevant sections fro indicate direct quotes from your mar information not in the ms, or briefly e	m manuso iuscript), o	cript title (i or elaborat	e on this it	tem by pro	viding add	litional		
"postoperative opioid utilization	"postoperative opioid utilization in orthopaedic trauma patients"							
1b) ABSTRACT: Structured s	summar	v of tria	al desiar	n. metho	ods. resi	ults. and		
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/functional		•			ention ar	nd		
comparator in the METHODS								
Mention key features/functionalities/components of the intervention and comparator in the abstract. If possible, also mention theories and principles used for designing the site. Keep in mind the needs of systematic reviewers and indexers by including important synonyms. (Note: Only report in the abstract what the main paper is reporting. If this information is missing from the main body of text, consider adding it)								
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subitem not at all important	0	0	0	<b>O</b>	0	essential		

Does your paper address su	bitem 1k	o-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 intervention group, who rece an ACT-based intervention for th received no messages"		-			_	_	
1b-ii) Level of human involve	ment in	the MF	THODS	section	of the A	RSTRACT	
Clarify the level of human involvement in the abstract, e.g., use phrases like "fully automated" vs. "therapist/nurse/care provider/physician-assisted" (mention number and expertise of providers involved, if any). (Note: Only report in the abstract what the main paper is reporting. If this information is missing from the main body of text, consider adding it)							
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Does your paper address su Copy and paste relevant sections fro this" to indicate direct quotes from y information not in the ms, or briefly e	m the mai	nuscript al script), or e	elaborate d	on this iter	n by provic	ling additional	
Your answer							

# 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

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

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subitem not at all important	0	0	0	0	•	essential

Does your paper	address subitem	1b-iv?
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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

"Eighty-two subjects were enrolled in the study. Seventy-six (38 ACT, 38 controls) completed the study. No differences between groups in demographic factors were identified. The intervention group utilized an average of 26.1±21.4 opioid tablets while the control group utilized 41.1±22.0 tablets, resulting in 36.5% less tablets utilized by subjects receiving the mobile phone-based ACT intervention (P= .004). Intervention group subjects reported a lower postoperative PROMIS Pain Intensity score of 45.9±7.2 compared to the control group's 49.7±8.8 (P= .04)."

# 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)

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# 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

Your answer

# INTRODUCTION

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

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

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

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# 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

"Public health concerns regarding opioid medications persist and healthcare systems are currently seeking solutions to the ongoing epidemic [1]."

"Acceptance and Commitment Therapy (ACT) is a specific behavior therapy that employs a pragmatic approach to help individuals decrease avoidable suffering and live according to self-identified personal values [11, 12]."

"Acceptance and Commitment Therapy has proven to be effective across multiple studies and patient populations in the treatment of pain [13]. Several studies report a high value for ACT in the management of chronic pain when compared to standard pharmacological treatment alone [13-16]."

"However, traditional ACT interventions have required a clinic based, interdisciplinary team approach, which is not always feasible for both patients and healthcare systems [17, 18]." "Evolving communication methods, such as automated mobile phone messaging [4, 19-21], for healthcare purposes are increasingly important as patients prefer these communication methods for delivering and receiving medical information [22]."

"Healthcare teams caring for patients with traumatic orthopaedic injuries have traditionally utilized opioid medication in the post-operative setting and these patients are at risk for prolonged opioid utilization in the postoperative period. We theorized the combination of ACT delivered via automated mobile phone messaging may help to decrease pain and opioid utilization in the acute postoperative setting. The aim of this prospective randomized controlled trial was to evaluate the effectiveness of ACT delivered via an automated mobile messaging robot on (1) decreasing early postoperative opioid utilization and (2) pain-related patient reported outcomes (PROs) in the first two weeks following surgery for acute, traumatic orthopaedic injuries."

# 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 appropriate), 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.

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# Does your paper address subitem 2a-ii? \*

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

"Additionally, every week of continued opioid utilization represents an increased risk of eventual misuse by patients [3]. Previous studies have found orthopaedic trauma patients utilize a decreasing amount of opioid medication in the first two postoperative weeks with six to fifteen days being the optimal opioid utilization period [4, 5]. In line with these findings, previous studies have utilized a two-week postoperative period to assess opioid medication consumption in surgical patients [4]."

"[11]. Acceptance and Commitment Therapy has proven to be effective across multiple studies and patient populations in the treatment of pain [13]. Several studies report a high value for ACT in the management of chronic pain when compared to standard pharmacological treatment alone [13-16]."

"Software driven, automated mobile phone messaging robots are low cost tools that can deliver predefined text-based information and receive incoming responses with high reliability when patients either prefer or it is necessary to communicate at distance [19, 23]. This technology demonstrates high efficacy as part of the treatment of conditions ranging from hypertension to substance abuse [24-26], and also has proved effective in increasing perioperative communication after hip and knee arthroplasty [27], and collecting pain and opioid medication data from patients following orthopaedic trauma and hand procedures [4, 19]."

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 prospective randomized controlled trial was to evaluate the effectiveness of ACT delivered via an automated mobile messaging robot on (1) decreasing early postoperative opioid utilization and (2) pain-related patient reported outcomes (PROs) in the first two weeks following surgery for acute, traumatic orthopaedic injuries."

#### **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

"Participants were randomized into either the control or intervention group using a standard online random number generator with a range set from 1 to 10 and a 1:1 ratio by a research assistant. Due to the nature of this study, subjects and the enrolling research assistant were not blinded to the participant's study group following randomization"

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

"Adults presenting to a university hospital level 1 trauma center indicated for operative

fixation of a traumatic upper or lower fracture were considered for the study (Table 2). Exclusion criteria are listed in Table 3. Eligible patients were approached prior to surgery by a research assistant in a private room. Individuals not excluded by screening questions and interested in participating underwent the informed consent process (Table 3). "
"One-hundred and twenty-five individuals were approached regarding the study over the five-month enrollment period between February and June 2019. Two patients were excluded at this time as they were non-English-speaking, and an additional 24 were excluded due to not utilizing mobile phone messaging or not having a personal mobile phone. This resulted in a total of 99 eligible people who were presented the study, 17 of which declined participation (Figure 1). Eighty-two subjects were enrolled, and six dropped from the study after providing consent due to various issues: one lost to follow-up, one withdrew at follow-up, one had incomplete follow-up, one operative plan changed to arthroplasty, and two subjects remained inpatient for over 7 days of the study period (Figure 1). "

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

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

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#### Does your paper address subitem 3b-i?

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

Your answer

# 4a) Eligibility criteria for participants

# Does your paper address CONSORT subitem 4a? \*

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

"Adults presenting to a university hospital level 1 trauma center indicated for operative fixation of a traumatic upper or lower fracture were considered for the study (Table 2). Exclusion criteria are listed in Table 3. Eligible patients were approached prior to surgery by a research assistant in a private room. Individuals not excluded by screening questions and interested in participating underwent the informed consent process (Table 3)."

# 4a-i) Computer / Internet literacy

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

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### Does your paper address subitem 4a-i?

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

Your answer

4a-ii) Open vs. closed, web-	based v	/s. face-	to-face	assessr	nents:	
Open vs. closed, web-based vs. face (online vs. offline), e.g., from an open based trial, or there were face-to-fac what degree got the study team to ki quasi-anonymous and whether havin measures (e.g., cookies, email confin	n access vecompone now the particular of the par	vebsite or ents (as pa articipant. identities	from a clir art of the i In online-o was possi	nic, and cla nterventio only trials, ible or whe	arify if this n or for ass clarify if pa ether techn	was a purely web- sessment), i.e., to articipants were lical or logistical
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4a-iii) Information giving duration giving duration given during recruitment informed consent procedures (e.g., pitem X26), as this information may his bias results.	. Specify loublish the	now partic informed	ipants wer consent d	ocumenta	tion as app	oendix, see also
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4b) Settings and locations v	vhere th	ne data	were co	ollected		
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4b-i) Report if outcomes we Clearly report if outcomes were (self-trials) or otherwise.				•		
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"At the time of consent subjects demographics questionnaire and Outcome Measure Information S Intensity 3A Short form, PROMIS Distress- Anxiety 8A Short form."	d baselin System (F Pain Int	e PROs c PROMIS)	onsisting Pain Inte	of the P	atient-Re <sub>l</sub> Short fori	ported m, PROMIS Pain
4b-ii) Report how institutions	al affilia	tions or	a dicalar	rod		
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5) The interventions for each group with sufficient details to allow replication, including how and when they were actually administered

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OWNERS  Mention names, credential, affiliationare owners or developer of the softwarentioned elsewhere in the manusc	vare, this n					
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5-ii) Describe the history/de Describe the history/development pr focus groups, usability testing), as the interpreting results.	rocess of t	he applica	tion and p			
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Revisions and updating. Clearly ment (and comparator, if applicable) evalua- during the evaluation process, or who Describe dynamic components such the replicability of the intervention (for	ated, or de ether the c as news f	escribe wh developme eeds or ch	ether the int and/or langing co	interventic content w intent whic	on underwe as "frozen"	ent major changes during the trial.
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5-vii) Access						
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# Does your paper address subitem 5-viii? \*

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

"The intervention group received twice-daily, text based mobile messages communicating an ACT-based intervention for the first two weeks following surgery (Multimedia Appendix 2). Control group subjects did not receive the ACT intervention or any other form of mobile message communications. The mobile messaging ACT protocol consisted of twice per day mobile messages, morning and evening, starting on postoperative day (POD) one and ending on POD 14. These mobile phone messages provided participants with an ACT-based intervention that was developed in collaboration with a pain psychologist (VK) specializing in ACT for chronic pain. These messages utilized all the principles presented in Table 1 with the objective of helping recipients understand and develop better coping skills in relation to their postoperative pain. An example message from Day 1 is below:

"Maintaining focus on what you value most in life is sometimes difficult after surgery. Do not let the momentary discomforts due to surgery take away from what you want most in life. Pick 3 things that matter most to you in life. Remind yourself of these 3 things you value most during your recovery process."

Outside of the mobile messaging intervention, both groups received the same standard postoperative care, healthcare team communications, and instructions for completing study follow-up. "

#### 5-ix) Describe use parameters

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

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

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

Your answer

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5-xi) Report any prompts/rer	nindors	usad				
Report any prompts/reminders used: use the application, what triggered the level of prompts/reminders required fapplication outside of a RCT setting (	Clarify if t em, freque or the tria	there were ency etc. I al, and the	t may be r level of pr	necessary ompts/rer	to distingu ninders fo	ish between the
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this is not applicable as the mobile phone messages themselves contained the intervention and did not require that patients access another application. No reminders for maintaining

pill counts were included in the study protocol.

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

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

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

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

This study had no co-interventions including training as the study only involved standard mobile phone messaging. Subjects falling under the categories of "No personal mobile phone with text messaging capabilities" or "Poor familiarity reading or sending mobile messages" were excluded from study participation.

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

"Baseline PROs were completed. Two weeks after operative intervention, follow-up was obtained in the form of an opioid medication pill count and postoperative administration of PROs. Mean number of opioid tablets utilized by patients were calculated and compared between groups. Mean PRO scores were also compared between groups."

"At the time of consent subjects were required to complete paper forms comprising a basic demographics questionnaire and baseline PROs consisting of the Patient-Reported Outcome Measure Information System (PROMIS) Pain Intensity 1A Short form, PROMIS Pain Intensity 3A Short form, PROMIS Pain Interference 8A Short form, and PROMIS Emotional Distress- Anxiety 8A Short form. Following completion of all PROs, participants were then randomized to their study group."

"The primary outcomes for this study was the amount of opioid pain medication consumed by subjects, and secondary outcomes analyzed were net change from baseline PRO scores at two-week follow-up."

"The method that participants employed to report their opioid medication consumption, and how PROs were captured during follow-up was recorded (Table 4). Subjects using their pill bottle to confirm the remaining number of opioid pain medication tablets from their discharge prescription on POD 14 were denoted as reporting a pill count. Cases where subjects or their care facility kept a log of tablet consumption were classified as reporting a daily log. Subjects reporting the number of tablets they used without the use of a log or pill count were designated as providing an estimate."

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

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

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Does your paper address subitem 6a-i?

Copy and paste relevant sections from manuscript text

Your answer

Describe whether and how "use" (incl (logins, logfile analysis, etc.). Use/ad- reported in any ehealth trial.						
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6a-iii) Describe whether, how was obtained						
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<b>7a) How sample size was de</b> NPT: When applicable, details of whe			ustering by	/ care prov	rides or cei	nters was
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7b) When applicable, explar	nation o	of any in	terim ar	nalyses	and stop	pping

# 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

This is not relevant to our study as no analysis were conducted outside of the two data collection points and we provided no specific stopping guidelines as participants could choose to remove consent at any time per our informed consent document.

### 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

"Participants were randomized into either the control or intervention group using a standard online random number generator with a range set from 1 to 10 and a 1:1 ratio by a research assistant. Due to the nature of this study, subjects and the enrolling research assistant were not blinded to the participant's study group following randomization."

# 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

"Participants were randomized into either the control or intervention group using a standard online random number generator with a range set from 1 to 10 and a 1:1 ratio by a research assistant. Due to the nature of this study, subjects and the enrolling research assistant were not blinded to the participant's study group following randomization."

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

"At the time of consent subjects were required to complete paper forms comprising a basic demographics questionnaire and baseline PROs consisting of the Patient-Reported Outcome Measure Information System (PROMIS) Pain Intensity 1A Short form, PROMIS Pain Intensity 3A Short form, PROMIS Pain Interference 8A Short form, and PROMIS Emotional Distress- Anxiety 8A Short form. Following completion of all PROs, participants were then randomized to their study group."

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 patients were approached prior to surgery by a research assistant in a private room. Individuals not excluded by screening questions and interested in participating underwent the informed consent process (Table 3). During consent, all subjects were informed of the outcomes of interests, different study arms, and that no changes would be made to their care in terms of postoperative medication, regardless of study participation."

"Participants were randomized into either the control or intervention group using a standard online random number generator with a range set from 1 to 10 and a 1:1 ratio by a research assistant."

"Subjects who received an odd number from the 1-10 range set on the random number generator were placed in the intervention group, while subjects given an even number were placed in the control group."

# 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 blinde	ed. and	who wa	sn't			
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11a-ii) Discuss e.g., whether pure "intervention of interest" and Informed consent procedures (4a-ii) of participants knew which intervention "comparator".	I which can create	one was	s the "co	mparat expectation	or" ons - discu	ss e.g., whether
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# 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

"During consent, all subjects were informed of the outcomes of interests, different study arms, and that no changes would be made to their care in terms of postoperative medication, regardless of study participation."

Participants were aware there was only one intervention.

# 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

This does not apply to this study as there is only a single intervention and subjects in the control group did not receive any mobile phone messages.

# 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

"To evaluate whether the intervention versus control group had a lower opioid utilization on average, we determined the number of tabs and MME taken in each group and compared means using t-tests."

"A separate power analysis was calculated for the PRO portion of the study and determined a total of 36 subjects would provide 80% power to detect a ten-point difference (1 standard deviation) in t-scores for the PROMIS instruments at an alpha level of .05. Statistical analyses were performed using SAS software version 9.4 (SAS Institute, Inc., Cary, NC)."

# 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]).

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

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

"Eighty-two subjects were enrolled, and six dropped from the study after providing consent due to various issues: one lost to follow-up, one withdrew at follow-up, one had incomplete follow-up, one operative plan changed to arthroplasty, and two subjects remained inpatient for over 7 days of the study period (Figure 1). This resulted in a final population 76 subjects (38 per study group). The enrollment period concluded once a powered sample for the primary aim was obtained. "

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

"Participant characteristics were described using mean± standard deviation (SD) or median (min-max) for continuous variables and frequencies (percentages) for categorical variables. Visual review of histograms and results of the Shapiro-Wilk test of continuous variables revealed only age and body mass index (BMI) were not normally distributed. Between group differences were evaluated using t-tests or Wilcoxon Rank Sum Tests (age and BMI) for continuous variables and chi-square or exact tests for categorical variables, as appropriate."

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

X26-i) Comment on ethics committee approval

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### 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 randomized controlled trial was registered with ClinicalTrials.gov (NCT03991546) and reporting is consistent with CONSORT guidelines (Multimedia Appendix 1). The study was performed at single center university hospital in Iowa City, IA, USA. Ethical approval of this study was provided by the University of Iowa Institutional Review Board and the study was determined Health Insurance Portability and Accountability Act compliant."

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#### **RESULTS**

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

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

#### Does your paper address CONSORT subitem 13a? \*

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

"One-hundred and twenty-five individuals were approached regarding the study over the five-month enrollment period between February and June 2019. Two patients were excluded at this time as they were non-English-speaking, and an additional 24 were excluded due to not utilizing mobile phone messaging or not having a personal mobile phone. This resulted in a total of 99 eligible people who were presented the study, 17 of which declined participation (Figure 1). Eighty-two subjects were enrolled, and six dropped from the study after providing consent due to various issues: one lost to follow-up, one withdrew at follow-up, one had incomplete follow-up, one operative plan changed to arthroplasty, and two subjects remained inpatient for over 7 days of the study period (Figure 1). This resulted in a final population 76 subjects (38 per study group)."

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

"One-hundred and twenty-five individuals were approached regarding the study over the five-month enrollment period between February and June 2019. Two patients were excluded at this time as they were non-English-speaking, and an additional 24 were excluded due to not utilizing mobile phone messaging or not having a personal mobile phone. This resulted in a total of 99 eligible people who were presented the study, 17 of which declined participation (Figure 1). Eighty-two subjects were enrolled, and six dropped from the study after providing consent due to various issues: one lost to follow-up, one withdrew at follow-up, one had incomplete follow-up, one operative plan changed to arthroplasty, and two subjects remained inpatient for over 7 days of the study period (Figure 1)."

#### 13b-i) Attrition diagram

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

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#### Does your paper address subitem 13b-i?

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

Your answer

14a) Dates defining the periods of recruitment and follow-up

#### Does your paper address CONSORT subitem 14a? \*

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

"One-hundred and twenty-five individuals were approached regarding the study over the fivemonth enrollment period between February and June 2019."

"The primary outcome for this study was the amount of opioid pain medication consumed by subjects, and secondary outcomes analyzed were net change from baseline PRO scores at two-week follow-up."

#### 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"

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#### 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

Your answer

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

#### Does your paper address CONSORT subitem 14b? \*

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

This trial was not stopped early. Enrollment for the trial was stopped once the powered sample of follow-ups had been achieved. The remaining few subjects without completed follow-up at that time were still utilized in the analysis, which is why we had 76 subjects instead of the powered sample of 74 needed.

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

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

#### Does your paper address CONSORT subitem 15? \*

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

These are all provided in table format within the manuscript.

"Table 4: Comparison of subject demographics by enrolled study group"

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

In ehealth trials it is particularly important to report demographics associated with digital divide issues, such as age, education, gender, social-economic status, computer/Internet/ehealth literacy of the participants, if known.

1

subitem not at all important

 $\bigcirc$   $\bigcirc$   $\bigcirc$ 

essential

#### Does your paper address subitem 15-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 trial utilized subjects who were familiar and comfortable with mobile phone messaging. No further digital divide issues were assessed in terms of internet or computer literacy. Demographics such as age and sex were included in the analysis.

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 "denom	16-i) Report multiple "denominators" and provide definitions					
Report multiple "denominators" and p study participation [and use] threshol used more than y weeks, N participan points of interest (in absolute and rela intervention.	ds" [1], e. ts "used"	g., N expo	sed, N con ention/cor	sented, N nparator a	used more t specific	e than x times, N pre-defined time
	1	2	3	4	5	
subitem not at all important	0	0	•	0	0	essential
Does your paper address sub	nitem 16	5-i2 *				
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Areas where multiple denominate contained footnotes to clarify if tused.  Table 4 footnote: "Data calculate All available data collected was ufrom patients who were determinate."	he enrol d using t utilized fo	led or fin final stud or analys	al study <sub>l</sub> y popula is, excep	oopulation tion only t in the c	n denom (N=38)."	inator was
16-ii) Primary analysis should Primary analysis should be intent-to-t the appropriate caveats that this is no	reat, seco longer a	ondary ana randomiz	lyses coul ed sample	(see 18-i)		only "users", with
	1	2	3	4	5	
subitem not at all important	0	0	0	0	0	essential
Does your paper address sub Copy and paste relevant sections fror indicate direct quotes from your mand information not in the ms, or briefly ex	n the mar uscript), o	nuscript (ir or elaborat	e on this i	tem by pro	viding add	itional

Your answer

# 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

"Utilizing a previous study of opioid medication usage in orthopaedic trauma patients [4], the sample size estimated to observe a 30% decrease in opioid utilization among two groups required a total of 74 subjects to achieve 80% power at an alpha of .05. "

"A separate power analysis was calculated for the PRO portion of the study and determined a total of 36 subjects would provide 80% power to detect a ten-point difference (1 standard deviation) in t-scores for the PROMIS instruments at an alpha level of .05. "

The precision of the effect size and its validity is based on the power analysis. This intervention is an adjunct to operative treatment.

# 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	0	0	0	0	0	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

Your answer

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

Does your paper address CC Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e No binary outcomes were preser	n the mar uscript), c xplain wh	nuscript (ii or elaborat y the item	nclude quo e on this it	em by pro	viding add	itional
18) Results of any other ana adjusted analyses, distingui				•	•	analyses and
Does your paper address CC Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e	n the mar uscript), c xplain wh	nuscript (ii or elaborat y the item	nclude quo e on this it is not app	em by pro licable/rel	viding add	itional
18-i) Subgroup analysis of co A subgroup analysis of comparing on stressed that this is a self-selected so (see 16-iii).	ly users is	s not unco	mmon in e			
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19) All important harms or (for specific guidance see CONSOR			cts in ea	ach grou	ıρ	
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All effects observed by the rese by study participants were repo breaches were reported during	rted in th	e study. N	-			
19-i) Include privacy breach		•				
Include privacy breaches, technical but also incidents such as perceived unexpected/unintended incidents. "I	d or real pr	ivacy breac	hes [1], te	chnical pro	oblems, an	d other
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No privacy breaches or technical problems were observed during this study.						

19-ii) Include qualitative feed staff/researchers	19-ii) Include qualitative feedback from participants or observations from staff/researchers					s from
Include qualitative feedback from pa strengths and shortcomings of the a or uses. This includes (if available) re by the developers.	pplication	, especiall	y if they po	oint to unii	ntended/ur	nexpected effects
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DISCUSSION						
22) Interpretation consister			balancir	ng bene	fits and	harms, and
considering other relevant NPT: In addition, take into account the			parator, la	ick of or p	artial blind	ing, and unequal
expertise of care providers or center						
22-i) Restate study question	s and su	ımmariz	e the ar	nswers s	uggeste	ed by the data,
starting with primary outcor	nes and	proces	s outcor	nes (us	e)	
Restate study questions and summa outcomes and process outcomes (us		nswers suç	ggested by	the data,	starting wi	ith primary
	1	2	3	4	5	

#### 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

"This randomized trial delivered an ACT-based intervention via an automated mobile messaging robot to post-operative orthopaedic patients. The subjects who received the ACT-based mobile phone intervention utilized a lower number of opioid tablets and consumed less MME in the first two weeks after their injury. We also found the intervention group reported less pain intensity and pain interference at two-week follow-up. This data demonstrates that ACT-based automated mobile messaging protocols may be effective in reducing the amount of opioid medication utilized and may positively affect postoperative PROs in patients undergoing operative fixation of their acute fractures."

"In this study we utilized ACT and a mobile phone messaging robot to assess whether these tools in combination could decrease opioid utilization and improve individual's perception of their early recovery from injury. Prior work has demonstrated a quicker time to opioid cessation and a decrease in postoperative opioid utilization (14% less in the ACT group) when utilizing in office ACT based treatments [17, 18]. Subjects receiving the ACT intervention via automated mobile phone message reported over 36% less opioid tablets and more than 34% less MME consumed than corresponding control subjects who did not receive ACT. Our findings suggest that software-based communication utilizing ACT through a mobile phone has the potential to have a large impact on utilization of postoperative pain medication by patients in the first weeks after surgery for fractures. Further study is required to determine if these effects are long lasting and to determine which injuries and patients get the greatest benefit. "

22-ii) Highlight unanswered n	•			: future ı	esearch	١
	1	2	3	4	5	
subitem not at all important	0	0	0	•	0	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

"Further study is required to determine if these effects are long lasting and to determine which injuries and patients get the greatest benefit. In addition, future investigations and trials should consider the effect of software delivery of ACT and other behavioral therapies on different cohorts of patients."

"Future research efforts may benefit from employing alternative PRO measures to identify the effects of ACT-based interventions including assessment of psychologic flexibility. Future research may also consider possible modification of our study protocol to include a longer intervention period and more than one follow-up data point. Future work may also consider designing an ACT based tool that is more focused on demonstrating an effect on PROs."

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 O O O o 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

"Several limitations were present in this study. First, we were limited to a single Level 1 trauma center, which may affect the reproducibility of our results across other healthcare settings. Next, the exclusion criteria for this study were extensive, and thus the results may not be generalizable to the entire scope of orthopaedic trauma patients. We attempted to include a diverse set of injuries and yet exclude patients with a high likelihood of confounding problems from open fractures or prolonged initial hospitalization. Future studies assessing the effects of ACT-based interventions like ours should aim for less restrictive exclusion criteria to apply this intervention to a larger, more diverse population. The research assistants were not blinded to the patient's study group. Additionally, patients understood the outcomes of interest in this study which could be susceptible to reporting bias. In addition, participants were not blinded to their treatment group."

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

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

#### 21-i) Generalizability to other populations

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

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

Does your p	paper address	subitem	21-i	?
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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

"Future studies assessing the effects of ACT-based interventions like ours should aim for less restrictive exclusion criteria to apply this intervention to a larger, more diverse population."

"First, we were limited to a single Level 1 trauma center, which may affect the reproducibility of our results across other healthcare settings."

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

Discuss if there were elements in the RCT that would be different in a routine application setting (e.g., prompts/reminders, more human involvement, training sessions or other co-interventions) and what impact the omission of these elements could have on use, adoption, or outcomes if the intervention is applied outside of a RCT setting.

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

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

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

Your answer

#### 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

"This randomized controlled trial was registered with ClinicalTrials.gov (NCT03991546) and reporting is consistent with CONSORT guidelines (Multimedia Appendix 1). "

### 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

"The intervention group received twice-daily, text based mobile messages communicating an ACT-based intervention for the first two weeks following surgery (Multimedia Appendix 2)."

# 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

"This study was possible thanks for a generous grant from the Orthopaedic Trauma Association."

#### X27) Conflicts of Interest (not a CONSORT item)

X27-i) State the relation of the In addition to the usual declaration of study team towards the system being identical with the developers/sponsor	f interests evaluate	s (financia ed, i.e., sta	l or otherw te if the au	rise), also	state the re	elation of the
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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 additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study  Your answer						
About the CONSORT EHEAL	.TH che	ecklist				
As a result of using this checomes, major changes yes, minor changes no	klist, di	d you m	ake cha	nges in	your ma	nuscript? *
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How much time did you spend on going through the checklist INCLUDING making changes in your manuscript *
3 hours
As a result of using this checklist, do you think your manuscript has improved? *
O yes
o no
Other:
Would you like to become involved in the CONSORT EHEALTH group?  This would involve for example becoming involved in participating in a workshop and writing an "Explanation and Elaboration" document
O yes
o no
Other:
Any other comments or questions on CONSORT EHEALTH
Your answer
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