# 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>

doi: 10.2196/jmir.1923

PMID: 22209829

,	Your name *
ı	First Last
	Cornelis Nils
	Primary Affiliation (short), City, Country * University of Toronto, Toronto, Canada
	KU Leuven, Leuven, Belgium
	Your e-mail address * abc@gmail.com
	nils.cornelis@kuleuven.be
	Title of your manuscript * Provide the (draft) title of your manuscript.
	Satisfaction and Acceptability of Telemonitored Home-Based Exercise in Patients With Intermittent Claudication: Pragmatic Observational Pilot 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.
(	Garmin Connect
	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")
Dutch
URL of your Intervention Website or App e.g. a direct link to the mobile app on app in appstore (itunes, Google Play), or URL of the website. If the intervention is a DVD or hardware, you can also link to an Amazon page.  https://connect.garmin.com/
URL of an image/screenshot (optional)  https://connect.garmin.com/
Accessibility * Can an enduser access the intervention presently?
<ul> <li>access is free and open</li> <li>access only for special usergroups, not open</li> <li>access is open to everyone, but requires payment/subscription/in-app purchases</li> <li>app/intervention no longer accessible</li> <li>Anders:</li> </ul>
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)"  General population

Primary Outcomes measured in trial * comma-separated list of primary outcomes reported in the trial
Satisfaction; acceptability
Secondary/other outcomes  Are there any other outcomes the intervention is expected to affect?
Walking capacity; physical fitness; quality of life; kinesiophobia; exercise self-efficacy; physical activity; adherence
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"
Anders:

Approx. Percentage of Users (starters) still using the app as recommended after 3 months $^{\ast}$
unknown / not evaluated
0-10%
O 11-20%
21-30%
31-40%
41-50%
51-60%
61-70%
71%-80%
81-90%
91-100%
Anders:
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
Anders: We found that patients were satisfied and accepted the use of a GPS

Article Preparation Status/Stage *  At which stage in your article preparation are you currently (at the time you fill in this form)
onot submitted yet - in early draft status
onot submitted yet - in late draft status, just before submission
submitted to a journal but not reviewed yet
submitted to a journal and after receiving initial reviewer comments
submitted to a journal and accepted, but not published yet
O published
Anders:
Journal *
If you already know where you will submit this paper (or if it is already submitted), please provide the journal name (if it is not JMIR, provide the journal name under "other")
onot submitted yet / unclear where I will submit this
Journal of Medical Internet Research (JMIR)
JMIR mHealth and UHealth
JMIR Serious Games
JMIR Mental Health
JMIR Public Health
JMIR Formative Research
Other JMIR sister journal
Anders:
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
Anders: 18739
TITLE AND ABSTRACT
1a) TITLE: Identification as a randomized trial in the title
1a) Does your paper address CONSORT item 1a? *  I.e does the title contain the phrase "Randomized Controlled Trial"? (if not, explain the reason under "other")
O yes
Anders: Our pilot is not a Randomized Controlled Trial.
1a-i) Identify the mode of delivery in the title  Identify the mode of delivery. Preferably use "web-based" and/or "mobile" and/or "electronic game" in the title. Avoid ambiguous terms like "online", "virtual", "interactive". Use "Internet-based" only if Intervention includes non-web-based Internet components (e.g. email), use "computer-based" or "electronic" only if offline products are used. Use "virtual" only in the context of "virtual reality" (3-D worlds). Use "online" only in the context of "online support groups". Complement or substitute product names with broader terms for the class of products (such as "mobile" or "smart phone" instead of "iphone"), especially if the application runs on different platforms.
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Does your paper address sub Copy and paste relevant sections from indicate direct quotes from your manu information not in the ms, or briefly ex	m manuso uscript), o	cript title (i or elaborate	e on this it	em by pro	viding add	itional
Satisfaction and Acceptability of Intermittent Claudication: Pragm					se in Pat	ients With
We opted for "Telemonitored" as using the online Garmin Connect received remotely feedback.	•	-		-		
1a-ii) Non-web-based components support").		•				
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1a-iii) Primary condition or ta Mention primary condition or target g Example: A Web-based and Mobile Int Randomized Controlled Trial	roup in th	e title, if a	ny (e.g., "fo			·
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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										
"Intermittent Claudication"										
1b) ABSTRACT: Structured s	summar	y of tria	al desigi	n, meth	ods, resi	ults, and				
NPT extension: Description of experi status.	mental tre	eatment, co	omparator	, care prov	iders, cent	ers, and blinding				
1b-i) Key features/functional	ities/coi	mponen	ts of the	e interve	ention ar	nd				
comparator in the METHODS	S sectio	n of the	ABSTRA	ACT						
Mention key features/functionalities/ possible, also mention theories and p systematic reviewers and indexers by what the main paper is reporting. If the adding it)	orinciples y including	used for d g importan	esigning t t synonym	he site. Ke is. (Note: '	eep in mind Only report	the needs of in the abstract				
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"The aim of this study is to asse HBET program using wearable to "Participants were instructed to exercise sessions per week in th	echnolog complete	y and ela e 3 walkir	stic band ng sessio	d resistar ons and 2	nce exerci elastic b	ses." AND and resistance				
In the results section of the abst highlight this part of the structur			he "app	oreciated	personal	feedback" to				

1b-ii) Level of human involved Clarify the level of human involveme "therapist/nurse/care provider/physif any). (Note: Only report in the absorbed from the main body of text, consider	nt in the al ician-assis ract what	ostract, e.g ted" (men the main p	g., use phra	ases like " er and exp	fully autom ertise of p	nated" vs. roviders involved,
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1b-iii) Open vs. closed, web- assessments in the METHOI Mention how participants were recru clinic or a closed online user group ( trial, or there were face-to-face compoutcomes were self-assessed through traditional offline trials, an open trial	OS section in the sec	on of the evs. offling ergroup tries part of the naires (as	e ABSTR ne), e.g., fro al), and cla he interver s common	ACT om an ope arify if this ntion or for in web-ba	en access v was a pur assessme sed trials)	website or from a ely web-based ent). Clearly say if . Note: In
researchers and participants know we "blinded" or "unblinded" to indicated usually refers to "open access" (i.e. the main paper is reporting. If this in	which treato the level o participant	ment is be of blinding s can self	eing admin instead of -enrol). (No	istered. To "open", as ote: Only re	avoid con "open" in eport in the	nfusion, use web-based trials e abstract what
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# Does your paper address subitem 1b-iii?

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

Not applicable for our study. From the abstract it is clear patients were engaging in a 4-week exercise program in their home-environment. In the methods section we elaborate on how recruitment was done (hospital based). During the exercise program all contact was limited to e-mail or telephone.

# 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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# 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

We added this information in the results section of our manuscript: "These results were reflected in high adherence to the prescribed walking sessions (GPS and logbook combined=mean 89%, SD 25; GPS only=mean 86%, SD 28), with 75% (15/20) of the patients completing all prescribed walking sessions. In contrast, patients were less compliant with resistance training (mean 85%, SD 22; 56% (9/16) completed all prescribed sessions and 20% (4/20) of patients did not return their logbook)".

1b-v) CONCLUSIONS/DISCUS	SSION i	n abstra	ct for ne	egative	trials	
Conclusions/Discussions in abstract negative (primary outcome not chang results are attributable to lack of uptamain paper is reporting. If this inform	ed), and t ike and di	he interve scuss reas	ntion was sons. (Not	not used, e: Only rep	discuss whoort in the	nether negative abstract what the
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INTRODUCTION						
2a) In INTRODUCTION: Scien	ntific ba	ackgrou	ınd and	explana	ition of	rationale
2a-i) Problem and the type of Describe the problem and the type of intervention vs. incorporated in broad population? Goals of the intervention, complement other solutions? (Note: E	system/s er health e.g., bein	colution the care progr g more co	at is objec am? Inten st-effectiv	ded for a <sub>l</sub> e to other	oarticular p interventio	oatient ons, replace or
Describe the problem and the type of intervention vs. incorporated in broad population? Goals of the intervention,	system/s er health e.g., bein Details ab	colution the care progr ng more co out the int	at is object am? Inten est-effectivervention	ded for a pre to other are provid	particular	oatient ons, replace or

# 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

Our introduction is build upon the rationale that several barriers are present for implementation of evidence-based supervised exercise programs ("SET is not readily available in most European countries, with only 30% of vascular surgeons having direct access [9,10]. Furthermore, even when SET is available, patients' participation is low, mainly because of a lack of transportation and time [11,12]." Yet, Home-Based alternatives are not as effective. Therefore, we wanted to investigate whether current technologies are to be considered in designing future home-based trials. This from a patient perspective.

"As such, wearable technology might help to bridge the gap by preserving the patient-provider relationship and offering home-based structured exercise therapy of adequate intensity in a health care system under pressure [14]."

Addition of elastic band exercises was also discussed, where the pain-free stimulus was hypothesized to be a popular alternative to traditional walking exercise.

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

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

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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

The study is performed from a patients' perspective. "However, one needs to address the needs and interests of all stakeholders, including patients [21]. In this line, a previous cohort study from our group showed that 81% of patients owning a computer and telephone were interested in telecoaching [25]. In addition, most patients preferred home-based exercise [26], and physiotherapists showed utmost interest (89%) in GPS tracking to monitor these sessions [27]." Our aim was to investigate whether this objectified interest would also translate in satisfaction, acceptability in this observational pilot.

# 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

Objectives (Page 2): "In this exploratory, pragmatic observational pilot study, we primarily aimed to evaluate patient satisfaction and acceptability of a structured model of HBET using wearable technology during walking, in combination with home-based resistance exercises. In addition, we aimed to report on the adherence and potential effectiveness of this combined intervention on walking capacity, physical fitness, physical activity levels, and quality of life in the development of an HBET program for patients with IC."

### **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

Methods (Page 2): "We conducted a 4-week exploratory observational cohort study to assess patient satisfaction and acceptability of an experimental HBET program combining walking therapy with elastic band exercises."

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

Does your paper address CC	NSORT	subiter	n 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								
Not applicable for our study.								
3b-i) Bug fixes, Downtimes, (	Content	: Change	es					
Bug fixes, Downtimes, Content Change changes to methods therefore also in during the trial (e.g., major bug fixes funexpected events" that may have in failures/downtimes, etc. [2].	cludes im or change	portant ches in the fu	nanges ma nctionality	ide on the or conter	interventiont) (5-iii) ar	on or comparator nd other		
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We think this is important in descour study. We used the Garmin C from user problems (referring to	onnect p	olatform a	and techr	nical issu	es were r	• •		
4a) Eligibility criteria for par	ticipan	ts						

# 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

Methods (Page 3): "Eligibility criteria included patients presenting with LEAD (Ankle-Brachial Index [ABI] ≤0.9 and/or a 15% decrease in ABI after a maximal treadmill test) and new-onset or conservatively treated IC (Rutherford I-III). Patients were excluded if they (1) had already participated in a structured, regular exercise program (eg, weekly physiotherapy); (2) showed exercise-induced signs of myocardial ischemia or complex ventricular arrhythmias during maximal treadmill exercise; (3) did not receive medical clearance for exercise; or (4) did not have access to a computer or the internet."

# 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

Methods (Page 3): Patients were excluded when they "(4) did not have access to a computer or the internet."

# 4a-ii) Open vs. closed, web-based vs. face-to-face assessments: Open vs. closed, web-based vs. face-to-face assessments: Mention how participants were recruited (online vs. offline), e.g., from an open access website or from a clinic, and clarify if this was a purely webbased trial, or there were face-to-face components (as part of the intervention or for assessment), i.e., to what degree got the study team to know the participant. In online-only trials, clarify if participants were quasi-anonymous and whether having multiple identities was possible or whether technical or logistical measures (e.g., cookies, email confirmation, phone calls) were used to detect/prevent these. subitem not at all important essential Selectie wissen Does your paper address subitem 4a-ii? \* Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study Methods (Page 3): "Patients consulting the ambulatory vascular center of the University Hospitals Leuven (Leuven) between October 2017 and July 2018 were recruited by vascular surgeons." Patients were contacted face-by-face by a researcher. 4a-iii) Information giving during recruitment Information given during recruitment. Specify how participants were briefed for recruitment and in the informed consent procedures (e.g., publish the informed consent documentation as appendix, see also item X26), as this information may have an effect on user self-selection, user expectation and may also bias results. subitem not at all important essential Selectie wissen Does your paper address subitem 4a-iii? Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study We do not explicitly state this in our manuscript. Patients were contacted after screening for eligibility by the vascular surgeons during routine consultations. A researcher went through the informed consent and assessed patient interest to be included in the pilot project.

4b) Settings and locations w	vhere th	ne data	were co	llected		
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This was added after reviewing t consultation at the vascular cent up measurements at our researc	ter, partio	cipants w	ere invite	ed for bas	seline and	•
4b-i) Report if outcomes were (self-trials) or otherwise.				•	•	
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# subitem not at all important O essential Selectie wissen Does your paper address subitem 4b-i? \* Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study Not applicable for our study. 4b-ii) Report how institutional affiliations are displayed Report how institutional affiliations are displayed to potential participants [on ehealth media], as affiliations with prestigious hospitals or universities may affect volunteer rates, use, and reactions with regards to an intervention.(Not a required item – describe only if this may bias results)

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5) The interventions for eac including how and when the	•				to allov	v replication,
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Important, yet not applicable in o	our study	<i>'</i> .				

5-ii) Describe the history/dev Describe the history/development pro focus groups, usability testing), as the interpreting results.	cess of tl	he applica	tion and p			, -
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Does your paper address sub Copy and paste relevant sections from indicate direct quotes from your mand information not in the ms, or briefly ex The application (Garmin Connect group (eg, TRICH-study NCT0204 intervention was further developed Intermittent Claudication.	n the mar uscript), o xplain why ) has be 7942, St	nuscript (in or elaborat y the item en used tart2spor	e on this it is not app in differe t NCT022	tem by pro licable/rel nt studie 240147).	oviding add levant for y s from ou This spe	litional vour study ur research cific
5-iii) Revisions and updating Revisions and updating. Clearly menti (and comparator, if applicable) evalua during the evaluation process, or whe Describe dynamic components such a the replicability of the intervention (for	ted, or de ther the d as news fe	escribe wh levelopme eeds or ch	ether the i nt and/or o anging co	nterventio content wantent which	n underwe as "frozen"	ent major changes during the trial.
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# 5-viii) Mode of delivery, features/functionalities/components of the intervention and comparator, and the theoretical framework

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

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

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Methods (Page 3) and figure 1: "To guide the 4-week home-based exercise program, participants were offered an informative booklet, a self-developed DVD with demonstration of the resistance band exercises (Multimedia Appendix 1), and a Garmin Forerunner 210." With regards to the home-based exercise program, patients received timely feedback as described (Methods, Page 3): "According to their individual preferences, participants received feedback twice weekly to only once during the 4-week intervention period via telephone or email." In addition, we relied upon behavioral change principles resulting from the proper use of GPS sport watches and the online platform: "This was personalized during contact moments using subjective reflection from the patients, baseline treadmill tests, and GPS-derived data. As such, participants had the possibility to self-monitor their walking sessions, received timely feedback on their performance, and were provided with information on how to adapt their walking program [31]."

5-ix) Describe use parameter Describe use parameters (e.g., intendrecommendations were given to the uwas the intervention used ad libitum.	ed "doses		-		-	
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5-xi) Report any prompts/rer Report any prompts/reminders used: use the application, what triggered the level of prompts/reminders required application outside of a RCT setting	: Clarify if t nem, frequ for 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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6a) Completely defined pre-specified primary and secondary outcome

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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

Not applicable for our study (no randomisation).

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

Not applicable for our study (no randomisation).

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

Not applicable for our study (no randomisation).

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

Specify who was blinded, and who wa	en't Heur	ally in web	-hacad tri	ale it ie no	nt noccible	to blind the
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# 11b) If relevant, description of the similarity of interventions

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

Does your paper address CONSORT subitem 11b? *  Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study  Not applicable for our 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	
This was adressed under the paragraph Statistics in Methods: "Statistical analyses were performed using JASP 0.11.1 (University of Amsterdam), with pre-post parametric (paired two-tailed t test) and nonparametric equivalent (Wilcoxon signed-rank) tests. An alpha level of 5% (two-sided) was used for statistical significance."	
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

Attrition was described in outcome tables, yet, no method was used to deal with missing values. Our analyses were mainly descriptive in a small sample (N=20).

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

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# 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

This was described in figure 2 of our manuscript (Page 5), with additional description under Results, Data Collection. In case of missing values, we highlighted this in the outcome tables (Supplementary files, Figure 3).

# 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

We only had one exercise group in our design. Patients flow was described in figure 2 of our manuscript (Page 5), with additional description and reason for exclusions under Results, Data Collection.

# 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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Baseline data are described in ta participant.	ble S1 (N	Multimed	ia appen	dix 2) for	each ind	ividual
15-i) Papart damagraphics a	ccociat	ad with	digital d	ivido iss	1100	
15-i) Report demographics as In ehealth trials it is particularly impo such as age, education, gender, social participants, if known.	rtant to re	eport demo	ographics	associated	d with digi	
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We did not specifically assess ehealth literacy or social-economic status. We described demographics mainly from a clinical perspective.

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

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

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

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# Does your paper address subitem 16-i? \*

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

We provide numbers of included patients in the outcome tables. In the results section, we highlight both group absolute and relative numbers, example in results on adherence (Page 5): "These results were reflected in high adherence to the prescribed walking sessions (GPS and logbook combined=mean 89%, SD 25; GPS only=mean 86%, SD 28), with 75% (15/20) of the patients completing all prescribed walking sessions. In contrast, patients were less compliant with resistance training (mean 85%, SD 22; 56% (9/16) completed all prescribed sessions and 20% (4/20) of patients did not return their logbook) and did not prefer this exercise alternative over conventional walking therapy (mean 2.65, SD 0.8; median 3, range: 1-5)."

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

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

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We did not include an intention-to	o-treat a	nalysis in	our obse	ervationa	ll pilot stu	ıdy.
17a) For each primary and se estimated effect size and its		•			•	•
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Summary data was described usi range where applicable.	ing mea	n or medi	an and s	tandard	deviation	or interquartile
17a-i) Presentation of process	s outco	mes suc	ch as me	etrics of	use and	d intensity of
In addition to primary/secondary (clin metrics of use and intensity of use (do not only refer to metrics of attrition (1 metrics such as "average session leng metric like a "session" is defined (e.g.,	ose, expo 3-b) (ofte gth". Thes	sure) and en a binary se must be	their opera variable), accompa	ational de but also t nied by a	finitions is o more co technica <b>l</b> c	critical. This does ntinuous exposure lescription how a
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18) Results of any other ana adjusted analyses, distingui				•	•	analyses and
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17b) For binary outcomes, presentation of both absolute and relative effect

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19) All important harms or u			cts in e	ach gro	up	
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19-II) Include qualitative feedback from participants or observations from staff/researchers								
Include qualitative feedback from participants or observations from staff/researchers, if available, on strengths and shortcomings of the application, especially if they point to unintended/unexpected effects or uses. This includes (if available) reasons for why people did or did not use the application as intended by the developers.								
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DISCUSSION								
22) Interpretation consistent considering other relevant of NPT: In addition, take into account the expertise of care providers or centers	evidenc e choice d	<b>e</b> of the com						
22-i) Restate study questions starting with primary outcomes and process outcomes (us	nes and	proces	s outcor	mes (use	e)	·		
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#### 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

Study aims were restated (Page 6, Discussion): "This study evaluated the satisfaction, acceptability, adherence, and potential effectiveness of a novel home-based exercise intervention that combines resistance training and walking therapy using wearables to monitor and guide patients with IC." All results were summarized, put into perspective and linked to previous findings (Page 6, Discussion, first paragraph).

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

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## 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

Page 7, Discussion: "Therefore, future studies should investigate the add-on effect of direct supervision in home-based interventions to (1) evaluate patient perception and methods to implement resistance exercises and (2) reduce activity-related fear using behavioral change or educational interventions."

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: Palook at a multiplicity of outcomes, incintervention/usability issues, biases the	reasing ri	sk for a Ty	pe I error.	Discuss b	iases due	to non-use of the
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the following sentence in the Disc the generalizability of this pilot in should be interpreted as such." A one researcher providing feedbac construct assessment with regar technology implementation: we de compared to the prescribed prog GPS watches and limitations who	tervention fter which ck and events to be did not as ram. In a	on, which  ch we cite  valuating  havioura  ssess the  addition,	n was par e differer all outco l change e accurac we highli	t of devent limitationes, the methods by of the ghted the	loping a lons in ou lack of p etc. With uploaded	arger trial and r protocol (eg, esychosocial h regards to sessions
21) Generalisability (externa NPT: External validity of the trial finding providers or centers involved in the tri	ngs accoi		•			•
21-i) Generalizability to other Generalizability to other populations: population, outside of a RCT setting, a results for other organizations	 In particu	lar, discus	-	-	_	
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Does your paper address subitem 21-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								
Discussion, limitations (Page 7): "Further limitations include the generalizability of this pilot intervention, which was part of developing a larger trial and should be interpreted as such."								
21-ii) Discuss if there were eleroutine application setting	ements	in the R	CT that	would b	e differ	ent in a		
Discuss if there were elements in the prompts/reminders, more human involument the omission of these element applied outside of a RCT setting.	olvement,	training se	essions or	other co-ir	nterventior	ns) and what		
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Although not an RCT, this observ clinical practice.	ational p	ilot was (	designed	to be ea	sily imple	emented in		
OTHER INFORMATION								
23) Registration number and	d 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

Methods, Page 2: "The study was approved by the Ethical Committee of UZ (ethics approval number: S59686; Belgian registration:B322201630074) Leuven/KU Leuven (Leuven, Belgium) and registered on ClinicalTrials.gov (NCT04043546)."

## 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

Methods, Page 2: "The study was approved by the Ethical Committee of UZ (ethics approval number: S59686; Belgian registration:B322201630074) Leuven/KU Leuven (Leuven, Belgium) and registered on ClinicalTrials.gov (NCT04043546)."

# 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

Not applicable for our study.

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

In addition to the usual declaration of the study team towards the system being evaluated In addition to the usual declaration of interests (financial or otherwise), also state the relation of the study team towards the system being evaluated, i.e., state if the authors/evaluators are distinct from or identical with the developers/sponsors of the intervention.								
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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  No conflict of interest.								
About the CONSORT EHEALTH checklist								
As a result of using this chec yes, major changes yes, minor changes no	klist, did	d you m	ake char	nges in Y	your ma	nuscript? *		
What were the most importa checklist?	nt chan	iges you	ı made a	as a resu	ılt of usi	ng this		
The checklist was used during re	vision of	f the enti	re manus	cript afte	er initial s	ubmission.		

How much time did you spend on going through the checklist INCLUDING making changes in your manuscript *
Approximately 240 minutes.
As a result of using this checklist, do you think your manuscript has improved? *
yes
O no
Anders:
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
Anders:
Selectie wissen
Any other comments or questions on CONSORT EHEALTH
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