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 (nonpharmacologic 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

Eysenbach G, CONSORT-EHEALTH Group

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

J Med Internet Res 2011;13(4):e126 URL: http://www.jmir.org/2011/4/e126/

doi: 10.2196/jmir.1923 PMID: 22209829

plunde@oslomet.no Bytt konto

Ikke delt

Utkastet er lagret

* indikerer at spørsmålet er obligatorisk

Your name * First Last Pernille Lunde Primary Affiliation (short), City, Country * University of Toronto, Toronto, Canada Oslo Metropolitan University, Oslo, Norway Your e-mail address * abc@gmail.com plunde@oslomet.no Title of your manuscript * Provide the (draft) title of your manuscript. Effects of individualized follow-up with an app post-cardiac rehabilitation: Five-years followup of a randomized controlled 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. Individualized follow-up with an app Evaluated Version (if any) e.g. "V1", "Release 2017-03-01", "Version 2.0.27913" Svaret ditt Language(s) * What language is the intervention/app in? If multiple languages are available, separate by comma (e.g. "English, French")

Norwegian

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. Svaret ditt URL of an image/screenshot (optional) Svaret ditt 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 The app used in the study is not accessible any more. Based on our fi Andre: 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)" Cardiovascular disease, noncommunicable dis Primary Outcomes measured in trial * comma-separated list of primary outcomes reported in the trial Peak Oxygen uptake Secondary/other outcomes Are there any other outcomes the intervention is expected to affect? Exercise performance, bodyweight, blood pressure, lipid profile, exercise habits, health related quality of life, health status, cardiac events, and physical activity

Recommended "Dose" * What do the instructions for users say on how often the app should be used?
Approximately Daily
Approximately Weekly
Approximately Monthly
Approximately Yearly
o "as needed"
Andre:
Approx. Percentage of Users (starters) still using the app as recommended after * 3 months
unknown / not evaluated
O-10%
O 11-20%
O 21-30%
O 31-40%
O 41-50%
51-60%
O 61-70%
71%-80%
O 81-90%
91-100%
Andre:

Overa	all, was the app/intervention effective? *
0)	yes: all primary outcomes were significantly better in intervention group vs control
()	partly: SOME primary outcomes were significantly better in intervention group vs control
O 1	no statistically significant difference between control and intervention
() .	potentially harmful: control was significantly better than intervention in one or more outcomes
O i	nconclusive: more research is needed
• P	Andre: At one-year follow-up the intervention was effective. However, as repo
	le Preparation Status/Stage * nich stage in your article preparation are you currently (at the time you fill in this form)
O 1	not submitted yet - in early draft status
O 1	not submitted yet - in late draft status, just before submission
O 5	submitted to a journal but not reviewed yet
0 9	submitted to a journal and after receiving initial reviewer comments
0 :	submitted to a journal and accepted, but not published yet
O 1	published
0 /	Andre:
-	nal * u already know where you will submit this paper (or if it is already submitted), please de the journal name (if it is not JMIR, provide the journal name under "other")
O 1	not submitted yet / unclear where I will submit this
O .	Journal of Medical Internet Research (JMIR)
0	JMIR mHealth and UHealth
0	JMIR Serious Games
0	JMIR Mental Health
0	JMIR Public Health
0	JMIR Formative Research
0	Other JMIR sister journal
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Andre: 60256						
TITLE AND ABSTRACT						
1a) TITLE: Identification as a	random	ized tria	l in the t	itle		
1a) Does your paper address I.e does the title contain the phra reason under "other")				ed Trial"?	' (if not, e	explain the
yes						
Andre:						
1a-i) Identify the mode of deli Identify the mode of delivery. Pro "electronic game" in the title. Avenue "Internet-based" only if Internemail), use "computer-based" or only in the context of "virtual reasupport groups". Complement of class of products (such as "mobapplication runs on different plants.	eferably oid amb vention "electro lity" (3-E r substit	use "web iguous te includes nic" only worlds) cute produ	rms like non-web if offline . Use "on uct name	"online", -based li products line" only es with b	"virtual", nternet co s are use v in the co roader te	"interactive". components (e.g. d. Use "virtual" context of "online rms for the
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subitem not at all important	0	0	•	0	0	essential
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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 The title includes "with an app" 1a-ii) Non-web-based components or important co-interventions in title Mention non-web-based components or important co-interventions in title, if any (e.g., "with telephone support"). 1 2 3 5 subitem not at all important essential Opphev valget Does your paper address subitem 1a-ii? Copy and paste relevant sections from manuscript title (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study This subitem seems not applicable 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 5 3 essential subitem not at all important Opphev valget Does your paper address subitem 1a-iii? * Copy and paste relevant sections from manuscript title (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study "post-cardiac rehabilitation"

Does your paper address subitem 1a-i? *

conclusions NPT extension: Description of ex and blinding status.	perimer	ntal treat	ment, co	mparato	r, care pro	oviders, centers,
1b-i) Key features/functionalit in the METHODS section of th Mention key features/functionalithe abstract. If possible, also me Keep in mind the needs of system synonyms. (Note: Only report in tinformation is missing from the results)	e ABST ties/cor ntion the natic re the abst	RACT nponent eories ar viewers a ract wha	s of the ind principend index and index t the mai	nterventi bles used kers by in in paper i	on and co I for design I cluding in I s reportir	omparator in gning the site. mportant
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1b-ii) Level of human involvem Clarify the level of human involved automated" vs. "therapist/nurse/ expertise of providers involved, it paper is reporting. If this informated adding it)	ement in care pro f any). (N	the abstovider/ph	tract, e.g. nysician-a y report	., use phr assisted" in the ab	ases like (mentior stract wh ly of text,	"fully n number and at the main

1b) ABSTRACT: Structured summary of trial design, methods, results, and

Does your paper address subitem 1b-ii?

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

The study of which this is a follow-up study of, is previously described in detail. Therefore, the human involvement is not the focus of this manuscript, and are reffered to in the main text. If a more comprehensive abstract is suggested from JMIR, we will rewrite the abstract accordingly.

1b-iii) Open vs. closed, web-based (self-assessment) vs. face-to-face assessments in the METHODS section of the ABSTRACT

Mention how participants were recruited (online vs. offline), e.g., from an open access website or from a clinic or a closed online user group (closed usergroup trial), and clarify if this was a purely web-based trial, or there were face-to-face components (as part of the intervention or for assessment). Clearly say if outcomes were self-assessed through questionnaires (as common in web-based trials). Note: In traditional offline trials, an open trial (open-label trial) is a type of clinical trial in which both the researchers and participants know which treatment is being administered. To avoid confusion, use "blinded" or "unblinded" to indicated the level of blinding instead of "open", as "open" in web-based trials usually refers to "open access" (i.e. participants can self-enrol). (Note: Only report in the abstract what the main paper is reporting. If this information is missing from the main body of text, consider adding it)

Does your paper address subitem 1b-iii?

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

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1b-iv) RESULTS section in abs Report number of participants e intervention (e.g., attrition/adher addition to primary/secondary of paper is reporting. If this information	nrolled/a rence me outcomes	assessed etrics, us s. (Note:	d in each e over tin Only repo	group, th ne, numb ort in the	per of log abstract	gins etc.), in t what the main
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1b-v) CONCLUSIONS/DISCUS Conclusions/Discussions in abs the trial is negative (primary out discuss whether negative result (Note: Only report in the abstract missing from the main body of t	tract for come no s are atti t what th	negative ot change ributable ne main p	e trials: Di ed), and th to lack o paper is r	iscuss the he interv	ne primar ention w and disc	as not used, cuss reasons.
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INTRODUCTION						

2a) In INTRODUCTION: Scient	tific bac	kground	l and exp	olanatio	n of ratio	onale
2a-i) Problem and the type of Describe the problem and the ty as stand-alone intervention vs. i particular patient population? Gother interventions, replace or cointervention are provided in "Me	pe of sys ncorpora oals of the	stem/sol ated in br ne interve ent other	ution tha oader he ention, e.	alth care g., being	program more co	n? Intended for a st-effective to
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2a-ii) Scientific background, r Scientific background, rationale: object of the study (be sure to d conditions/diagnoses, if approp and what is the context for this study performed, potential impa comparator.	: What is iscuss th iate), mo specific	known a ne use of otivation study, fro	bout the similar s for the st m which	(type of) systems f udy, i.e. v stakeho	system or other vhat are lder view	that is the the reasons for point is the
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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

"In 2020, we demonstrated that individualized follow-up for one year with an app improved adherence to healthy behaviour post-CR. In this study, 113 patients were randomly assigned to individualized follow-up enabled with an app, or to a control group (CG) which received usual care. At one-year follow-up there were a statistically significant difference between the groups, in favour to the intervention group (IG), in V02peak (ml/kg/min), exercise performance, exercise habits and in self-perceived goal achievement. A unique finding was the low dropout rate (n=2) as well as the high levels of adherence to the app. As far as the authors know, our study was the first evaluating the effect of an app for a whole year in a post-CR setting. To date, however, there is no research investigating the long-term effects of successful mHealth interventions beyond one year when the use of app discontinues."

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 primary aim of this study was to examine whether the effect demonstrated in VO2peak after one year was maintained four years post intervention. In the same manner, as secondary outcomes, we aimed to evaluate exercise performance, bodyweight, resting blood pressure (BP), lipid profile, exercise habits, health-related quality of life (HRQL), health status, cardiac events, and level of physical activity.

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

"We conducted a single-blinded, RCT with two arms. Patients were allocated to a CG or to an IG in a 1:1 ratio. A computer-generated, permuted block randomization scheme was used."

3b) Important changes to me criteria), with reasons	thods a	ifter trial	comme	ncemen	t (such a	as eligibility
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As this is a follow-up study of a I design are reported in the feasib					-	
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Bug fixes, Downtimes, Content C description of changes to metho the intervention or comparator of functionality or content) (5-iii) ar study design such as staff chan-	Changes ods there during th nd other	efore also efore trial (e. "unexpe	systems o include: g., major cted ever	s importa bug fixe: nts" that	ant chang s or chan may have	ges made on ges in the
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4a) Eligibility criteria for partic	cipants					

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

Like this: "inclusion criteria were patients completing one of the CR programs, age ≥40 years, owner and user of an Android or Apple smartphone, and able to read and understand Norwegian or English language. Exclusion criteria included ischemia or arrhythmias uncovered at cardiopulmonary exercise test (CPET) that gave restrictions equivalent to <80% of maximal heart rate or BORG scale (6-20) <15 at exercise. Further, patients with muscular or skeletal disorders that affected exercise capacity more than the heart disease were excluded. Furthermore, patients with severe malignant disease (i.e., advanced cancer) that affected the patient`s life span to a greater extent than their heart disease were also excluded."

4a-i) Computer / Internet liter	acy					
Computer / Internet literacy is o explicitly clarified.	often an ir	mplicit "d	le facto"	eligibility	criterion	- this should be
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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

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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 web-based 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.

cookies, email confirmation, pho	one calls) were us	sed to de	tect/prev	ent thes	e.
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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

Patients were recruited offline: "Patients were recruited from two CR centres in the eastern part of Norway. Whereas one of the centres offer one- and four-week inpatient CR programs, and the other centre offers a 12-weeks outpatient CR program."

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.

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Does your paper address subitem 4a-iii?

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It is previously described in detail how the recruitment were handled (Lunde et al 2020)

4b) Settings and locations where the data were collected

Does your paper address CONSORT subitem 4b? *

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

"Patients were recruited from two CR centres in the eastern part of Norway. Whereas one of the centres offer one- and four-week inpatient CR programs, and the other centre offers a 12-weeks outpatient CR program."

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including how and when they were actually administered

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5-iii) Revisions and updating Revisions and updating. Clearly application/intervention (and co- intervention underwent major or development and/or content was such as news feeds or changin the intervention (for unexpected	omparato hanges d as "frozer g content	or, if appli during the n" during t which m	cable) ev evaluati the trial. nay have	raluated, on proce Describe	or descri ess, or wh e dynamic	be whether the ether the components
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5-iv) Quality assurance meth	ods					
Provide information on quality a information provided [1], if appl	assuranc	e method	ls to ens	ure accu	racy and	quality of
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5-v) Ensure replicability by publishing the source code, and/or providing screenshots/screen-capture video, and/or providing flowcharts of the algorithms used Ensure replicability by publishing the source code, and/or providing screenshots/screen-capture video, and/or providing flowcharts of the algorithms used. Replicability (i.e., other researchers should in principle be able to replicate the study) is a hallmark of scientific reporting.									
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5-vii) Access Access: Describe how participants accessed the application, in what setting/context, if they had to pay (or were paid) or not, whether they had to be a member of specific group. If known, describe how participants obtained "access to the platform and Internet" [1]. To ensure access for editors/reviewers/readers, consider to provide a "backdoor" login account or demo mode for reviewers/readers to explore the application (also important for archiving purposes, see vi).									
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subitem not at all important	0	0	0	•	0	essential			
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Does your paper address subitem 5-vii? * Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study "After baseline assessment, patients who were randomly assigned to the IG were given access to an app and received training on its use."									
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? *

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

In addition to the previously published papers on the intervention (JMIR formative research, BMC Sports Science, Medicine and Rehabilitation and European journal of preventive cardiology), following has been included in the submitted manuscript: "Intervention group

After baseline assessment, patients who were randomly assigned to the IG were given access to an app and received training on its use. The app was personalized with the patients' goals and tasks for the upcoming year. Each patient decided when and how often they wanted reminders of their tasks. They could also use the app for reflections or notes related to a specific goal and directly communicate any questions to the supervisor. The supervisor (specialized physiotherapist) had access to an administrator interface and monitored each patient. The interface also allowed the supervisor to send short motivational messages (maximum 112 characters) directly to each patient through the app. Patients in the IG received comprehensive individualized feedback based on their completed tasks, pending tasks and/or any notes they had written. This feedback was provided weekly the first 12 weeks and monthly for the rest of the year. Additionally, all patients received one to three short individualized motivational messages per week throughout the year. Questions submitted to the supervisor were answered within two working days. This individualized follow-up continued until one-year follow-up assessment. A detailed description of the intervention is presented elsewhere [10, 11]. After the one-year follow-up, the app was removed from the patient's smartphones and the follow-up process were discontinued. Patients were encouraged to maintain or improve their healthy behaviours based on their individual risk profiles.

Control group

Patients allocated to the CG received treatment as usual. This included visiting their general practitioner and cardiologist, if, and when needed. After baseline assessment, they were encouraged to maintain or improve healthy behaviour based on their needs and goals. Encouragement to maintain or improve healthy behaviour was given at the one-year follow-up assessment. "

5-ix) Describe use parameter	S							
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 sub	item 5-i	x?								
Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study										
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5-x) Clarify the level of human Clarify the level of human involved technical assistance) in the e-inexpertise of professionals involued and frequency of the support, his delivered". It may be necessarequired for the trial, and the level outside of a RCT setting (discussion).	vement (contervention ved, if an ow it is in the content of the content velocities where the content velocities wellong to the content velocities where the content velocities wellong the content velocities where the content velocities wellong the content velocities where velocities were velocitie	care prov on or as c y, as well nitiated, a inguish b nan invol	o-interve l as "type and the m between t vement r	ntion (de of assis ledium b he level equired f	tail numl tance off y which t of humar or a rout	oer and ered, the timing he assistance n involvement				
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5-xi) Report any prompts/reminders used Report any prompts/reminders used: Clarify if there were prompts (letters, emails, phone calls, SMS) to use the application, what triggered them, frequency etc. It may be necessary to distinguish between the level of prompts/reminders required for the trial, and the level of prompts/reminders for a routine application outside of a RCT setting (discuss under item 21 – generalizability).										
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Does your paper address subitem 5-xi? *

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

This is described in the submitted manuscript, see 5-viii. In addition, it is even more elaborated in previously studies on the same intervention.

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 was a stand alone intervention. All components in the app has been described in detail.

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

Primary and secondary outcome measures are described in detail as well as how they were assessed. See the following paragraphs in the sumitted manuscript: "Outcomes and assessments" "Peak oxygen uptake and exercise performance" "Bodyweight", "Blood pressure" "Lipid profile" "Exercise Habits" "Health related quality of life" "Health status" "Cardiac events" and "Physical activity"

6a-i) Online questionnaires: d CHERRIES items to describe If outcomes were obtained thro for online use and apply CHERR designed/deployed [9].	how the	questio le questio	nnaires onnaires,	were de , describe	signed/o	deployed vere validated			
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Does your paper address subitem 6a-i? Copy and paste relevant sections from manuscript text Svaret ditt									
6a-ii) Describe whether and how "use" (including intensity of use/dosage) was defined/measured/monitored Describe whether and how "use" (including intensity of use/dosage) was defined/measured/monitored (logins, logfile analysis, etc.). Use/adoption metrics are important process outcomes that should be reported in any ehealth trial. 1 2 3 4 5									
Does your paper address subitem 6a-ii? Copy and paste relevant sections from manuscript text Svaret ditt									
6a-iii) Describe whether, how, and when qualitative feedback from participants was obtained Describe whether, how, and when qualitative feedback from participants was obtained (e.g., through emails, feedback forms, interviews, focus groups).									
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Does your paper address subitem 6a-iii?

Copy and paste relevant sections from manuscript text

Page 16: "This was also emphasized by the patients in a qualitative study that aimed to investigate experiences with the follow-up provided [29]. The patients emphasized that the person behind the app (the supervisor) was the most significant success factor in promoting adherence to healthy behaviour [29]. "

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

Does your paper address CONSORT subitem 6b? *

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

There were no changes in trial outcomes after the trial commenced

7a) How sample size was determined

NPT: When applicable, details of whether and how the clustering by care provides or centers was addressed

7a-i) Describe whether and how expected attrition was taken into account when calculating the sample size

Describe whether and how expected attrition was taken into account when calculating the sample size.

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Does your paper address subitem 7a-i?

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

"Sample size was calculated from the primary outcome, assuming a difference in relative VO2peak between groups of 3.5 ml/kg/min as a clinical important difference [23-25]. Based on a previously conducted feasibility study [26], we estimated the associated standard deviation (SD) to be 6 ml/kg/min. With a power of 80% and significance level of 0.05, the sample size was calculated to be sufficient with 47 patients in each group. To allow for a 20% dropout at one-year follow-up, we aimed to include 113 patients in total."

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

Does your paper address CONSORT subitem 7b? *

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

Unfortunately, I dont know the content of this question

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

"Patients were allocated to a CG or to an IG in a 1:1 ratio. A computer-generated, permuted block randomization scheme was used. The randomization was stratified by CR program and patients were randomly allocated to one of the two groups via concealed allocation right after baseline assessment."

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

"Patients were allocated to a CG or to an IG in a 1:1 ratio. A computer-generated, permuted block randomization scheme was used. The randomization was stratified by CR program and patients were randomly allocated to one of the two groups via concealed allocation right after baseline assessment."

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

Patients were allocated to a CG or to an IG in a 1:1 ratio. A computer-generated, permuted block randomization scheme was used. The randomization was stratified by CR program and patients were randomly allocated to one of the two groups via concealed allocation right after baseline assessment.

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

It is described in detail in the protocol paper, published in BMC Sports Science, Medicine and Rehabilitation. This paper has been cited in the submitted manuscript, but the prosess is also described in the methods section.

11a) If done, who was blinded after assignment to interventions (for example, participants, care providers, those assessing outcomes) and how NPT: Whether or not administering co-interventions were blinded to group assignment

11a-i) Specify who was blinded, and who wasn't

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

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Does your paper address subitem 11a-i? *

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

"Test personnel measuring the primary outcome were blinded for group allocation." In addition, objectively measured measurements, such as bodyweight, blood pressure and lipid profile, the measurement was blinded for group allocation.

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

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

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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

Svaret ditt

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

There were no similarities as the intervention group was compared with a control group whih received usual care (nothing)

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

"IBM SPSS Statistics (version 29) and STATA (version 18) were used for statistical analysis. Continuous, normally distributed baseline data were analysed with an independent t-test to test for differences between groups, and Pearson's chi-squared test was used to analyse the categorical data. Baseline differences between cases with and without five-year outcome data were analysed using the same statistical tests. The assessment of missing data was done based on strategies for dealing with missing data in clinical trials [27]. Differences between groups for primary and secondary outcomes were assessed using a mixed model for repeated measurements with a subject-specific random intercept and group, follow-up time, interaction between group and follow-up time, and the outcome measure at baseline as fixed effects. The Mann Whitney U test was applied to analyse for differences between groups in IPAQ at five-year follow-up. Analysis was carried out by intention-to-treat and all tests were two-sided. Data are presented as mean ± SD unless stated otherwise. A p-value <0.05 was considered significant."

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

"The assessment of missing data was done based on strategies for dealing with missing data in clinical trials [27]. " "There were no statistically significant differences in the baseline characteristics between cases with missing data at five-years follow-up and cases with no missing data. We assumed missing data as completely at random, since they were unrelated to any observed or unobserved variables [27]. Thus, imputation of missing data was not conducted [27]. "

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

Yes, the analyses is adjusted based on the baseline values. This is described by the coauthor Prof Pripp which is the statistician in our research group.

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

"The Regional Committee for Medical and Health Research Ethics (South-East ID: 2016-1476) approved the study protocol and the study was conducted in accordance with the Helsinki Declaration. For the present follow-up study, an updated study protocol was sent and approved. "

x26-ii) Outline informed consent procedures

Outline informed consent procedures e.g., if consent was obtained offline or online (how?

Checkbox, etc.?), and what information was provided (see 4a-ii). See [6] for some items to be included in informed consent documents.							
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Does your paper address subitem X26-ii? Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study Svaret ditt X26-iii) Safety and security procedures Safety and security procedures, incl. privacy considerations, and any steps taken to reduce the likelihood or detection of harm (e.g., education and training, availability of a hotline) 1 2 3 essential subitem not at all important Does your paper address subitem X26-iii? Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study Svaret ditt **RESULTS** 13a) For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analysed for the primary outcome NPT: The number of care providers or centers performing the intervention in each group and the number of patients treated by each care provider in each center Does your paper address CONSORT subitem 13a? * Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study A consort flow-chart is provided at page 10 in the submitted manuscript 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

This is clearly reported in the manuscript at page 10.

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

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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

"In total, 177 patients were screened for eligibility at two CR centres from October 2017 until June 2018 whereas 113 were included and randomized to the IG or CG (see Figure 1). Five-years follow-up were completed in September 2023, and in total, 101 patients were assessed."

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 sub	oitem 14	a-i?								
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Svaret ditt	Svaret ditt									
14b) Why the trial ended or was stopped (early)										
Does your paper address CO	Does your paper address CONSORT subitem 14b? *									
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THe trial did not end or stop ear	·ly									
15) A table showing baseline group NPT: When applicable, a descri										
expertise, etc.) and centers (vol					, 4					
Does vour naner address CO	NSORT (suhitem	152 *							
Copy and paste relevant section "like this" to indicate direct quo providing additional information	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									
THis is provided in table 1										

15-i) Report demographics as	sociate	d with d	igital div	ide issu	es		
In ehealth trials it is particularly i divide issues, such as age, educ computer/Internet/ehealth litera	ation, ge	nder, soc	ial-econ	omic sta		ed with digital	
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Does your paper address subi	item 15	-j? *					
Copy and paste relevant section: "like this" to indicate direct quot providing additional information applicable/relevant for your stud This is provided in table 1. In add	es from not in th	your mar ne ms, or	nuscript) briefly ex	, or elabo xplain wh	orate on t by the iter	his item by m is not	
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 "denomi	nators"	and prov	vide defi	nitions			
Report multiple "denominators" a "across a range of study particip consented, N used more than x t intervention/comparator at spec relative numbers per group). Alw	ation [ar times, N cific pre-c	nd use] th used mo defined ti	nresholds re than y me point	s" [1], e.g weeks, I ts of inte	., N expo N particip rest (in al	sed, N pants "used" the bsolute and	
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16-ii) Primary analysis should Primary analysis should be inter only "users", with the appropriat 18-i).	nt-to-trea	t, second	lary anal	•					
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Does your paper address subitem 16-ii? Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study "Analysis was carried out by intention-to-treat and all tests were two-sided."									
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 This is provided in tables and text in the result section.									
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).									
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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 Svaret ditt 17b) For binary outcomes, presentation of both absolute and relative effect sizes is recommended Does your paper address CONSORT subitem 17b? * Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study Reporting on all outcomes follows the prefered reporting based on the measurements development. 18) Results of any other analyses performed, including subgroup analyses and adjusted analyses, distinguishing pre-specified from exploratory Does your paper address CONSORT subitem 18? * Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study The submitted manuscript includes all analyses performed. Subgroup analyses has not been conducted. 18-i) Subgroup analysis of comparing only users A subgroup analysis of comparing only users is not uncommon in ehealth trials, but if done, it must be stressed that this is a self-selected sample and no longer an unbiased sample from a randomized trial (see 16-iii). 1 2 3 4 5 0 0 0 0 subitem not at all important essential

Does your paper address sub	oitem 18	-1?							
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									
Svaret ditt									
19) All important harms or unintended effects in each group (for specific guidance see CONSORT for harms)									
Does your paper address CO	NSORT s	subitem	19? *						
Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study									
There were no uninteded effects CONSORT flow-chart in addition			-	hing sho	uld be cle	ear based on the			
19-i) Include privacy breache Include privacy breaches, techn participants, but also incidents problems, and other unexpected unintended positive effects [2].	ical prob such as p	lems. Thi	s does n I or real p	rivacy b	reaches [1], technical			
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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.									
	1	2	3	4	5				
subitem not at all important	0	0	0	0	0	essential			
Does your paper address subitem 19-ii? Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study Svaret ditt									
DISCUSSION									
considering other relevant ev NPT: In addition, take into accor	22) Interpretation consistent with results, balancing benefits and harms, and considering other relevant evidence NPT: In addition, take into account the choice of the comparator, lack of or partial blinding, and unequal expertise of care providers or centers in each group								
starting with primary outcom Restate study questions and su	22-i) Restate study questions and summarize the answers suggested by the data, starting with primary outcomes and process outcomes (use) Restate study questions and summarize the answers suggested by the data, starting with primary outcomes and process outcomes (use).								
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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

"Our primary finding indicates that the beneficial effects observed in VO2peak and exercise performance one-year post-CR [10] have ceased away four-year post intervention. A strength of our study is the low drop-out rate at both one- and five-year follow up. For the primary outcome (VO2peak), we maintained adequate statistical power to detect a genuine difference between the groups. For secondary outcomes, the results must be interpreted with caution."

22-ii) Highlight unanswered n Highlight unanswered new ques	-				search	
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20) Trial limitations, addressi relevant, multiplicity of analys	-	ces of p	otential	bias, im	orecisio	n, and, if
20-i) Typical limitations in ehe Typical limitations in ehealth tria trials often look at a multiplicity biases due to non-use of the interconsent procedures, unexpected	als: Parti of outco ervention	cipants i omes, inc n/usabili	reasing r	isk for a	Type I eri	ror. Discuss
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This is not that relevant as it is a beforementioned studies of which		-		_			
21) Generalisability (external NPT: External validity of the trial patients, and care providers or c	findings	accordi	ng to the	intervent			
21-i) Generalizability to other Generalizability to other populati Internet population, outside of a applicability of the study results	ons: In p RCT set	oarticular ting, and	general	•	-	•	
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21-ii) Discuss if there were eleapplication setting Discuss if there were elements in setting (e.g., prompts/reminders interventions) and what impact to adoption, or outcomes if the interventions.	n the RC , more h .he omis	T that wo uman inv	ould be d olvemer nese eler	ifferent ir nt, training ments co	n a routing g session uld have	e application us or other co- on use,	
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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

Svaret ditt

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

"The original study protocol and the updated study protocol were registered in ClinicalTrials.gov with the following IDs NCT03174106 and NCT05697120 respectively."

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

ClinicalTrials.gov with the following ID NCT05697120. In addition, the protocol paper was published in 2019 in BMC Sports Science, Medicine and Rehabilitation.

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 project is funded by Foundation Dam (grant number 326568)."

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What were the most importal checklist?						
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As a result of using this checklist, do you think your manuscript has improved? *
yes
O no
Andre: The checklist seems good for eHealth studies. However, as the prese
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
yes
O no
Andre: Yes, If it is a clear mandat and workgroup.
Opphev valget
Any other comments or questions on CONSORT EHEALTH
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