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

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

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

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

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

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

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

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

Mandatory reporting items are marked with a red *.

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

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

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

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

Eysenbach G, CONSORT-EHEALTH Group

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

Mobile Health Interventions

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

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

doi: 10.2196/jmir.1923

PMID: 22209829



48553@phec.de (not shared) Switch account



Draft saved

* Required

Your name *

First Last

Jennifer Muschol

Primary Affiliation (short), City, Country *

University of Toronto, Toronto, Canada

Justus Liebig University Giessen, Giessen, Ger

Your e-mail address *

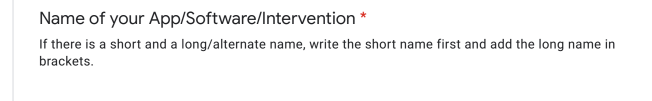
abc@gmail.com

Jennifer.Muschol@wirtschaft.uni-giessen.de

Title of your manuscript *

Provide the (draft) title of your manuscript.

Assessing Telemedicine Efficiency in German Follow-up Care With Video Consultations for Patients in Orthopedic and Trauma Surgery: A Randomized Controlled Trial



Evaluated Version (if any)

e.g. "V1", "Release 2017-03-01", "Version 2.0.27913"

Your answer

CLICKDOC

Language(s) *

What language is the intervention/app in? If multiple languages are available, separate by comma (e.g. "English, French")

German

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://clickdoc.elvi.de/#/login

URL of an image/screenshot (optional)

Your answer

Accessibility * Can an enduser access the intervention presently?
access is free and open
access only for special usergroups, not open
access is open to everyone, but requires payment/subscription/in-app purchases
app/intervention no longer accessible
Other:
Primary Medical Indication/Disease/Condition *
e.g. "Stress", "Diabetes", or define the target group in brackets after the condition, e.g. "Autism (Parents of children with)", "Alzheimers (Informal Caregivers of)"
Orthopedic and trauma surgery follow-up
Primary Outcomes measured in trial * comma-separated list of primary outcomes reported in the trial
communication of primary outcomes reported in the than
Patient satisfaction
Secondary/other outcomes
Are there any other outcomes the intervention is expected to affect?
Physician satisfaction, Quality of care

Recommended "Dose" * What do the instructions for users say on how often the app should be used?
Approximately Daily
Approximately Weekly
Approximately Monthly
Approximately Yearly
as needed"
Other: No App, One-time video consultation
Approx. Percentage of Users (starters) still using the app as recommended after 3 months *
unknown / not evaluated
0-10%
11-20%
21-30%
31-40%
41-50%
51-60%
61-70%
71%-80%
81-90%
91-100%
Other:

Overall, was the app/intervention effective? *
yes: all primary outcomes were significantly better in intervention group vs control
partly: SOME primary outcomes were significantly better in intervention group vs control
o no statistically significant difference between control and intervention
outcomes potentially harmful: control was significantly better than intervention in one or more
inconclusive: more research is needed
Other:
Article Preparation Status/Stage * At which stage in your article preparation are you currently (at the time you fill in this form)
At which stage in your article preparation are you currently (at the time you fill in this form)
At which stage in your article preparation are you currently (at the time you fill in this form) not submitted yet - in early draft status
At which stage in your article preparation are you currently (at the time you fill in this form) not submitted yet - in early draft status not submitted yet - in late draft status, just before submission
At which stage in your article preparation are you currently (at the time you fill in this form) not submitted yet - in early draft status not submitted yet - in late draft status, just before submission submitted to a journal but not reviewed yet
At which stage in your article preparation are you currently (at the time you fill in this form) not submitted yet - in early draft status not 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
At which stage in your article preparation are you currently (at the time you fill in this form) onot submitted yet - in early draft status not submitted yet - in late draft status, just before submission submitted to a journal but not reviewed yet submitted to a journal and after receiving initial reviewer comments submitted to a journal and accepted, but not published yet

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
Other:
Is this a full powered effectiveness trial or a pilot/feasibility trial? *
O 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)
o no ms number (yet) / not (yet) submitted to / published in JMIR
Other:

TITLE AND ABSTRACT						
1a) TITLE: Identification as a	randor	nized tr	ial in th	e title		
1a) Does your paper address I.e does the title contain the phrase "F "other") yes Other:				(if not, ex	plain the re	eason under
1a-i) Identify the mode of delivery. Preferal title. Avoid ambiguous terms like "onl includes non-web-based Internet comoffline products are used. Use "virtua only in the context of "online support terms for the class of products (such application runs on different platform	bly use "w line", "virtu nponents (Il" only in t groups". (as "mobi	veb-based' ual", "inter (e.g. email the contex Compleme	and/or "n active". Us), use "cor t of "virtua ent or subs	se "Interne mputer-bas al reality" (stitute pro	t-based" or sed" or "ele 3-D worlds duct names	nly if Intervention ectronic" only if). Use "online" s with broader
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Your answer						
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1b) ABSTRACT: Structured summary of trial design, methods, results, and conclusions

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

1b-i) Key features/functionalities/components of the intervention and comparator in the METHODS section of the ABSTRACT

Mention key features/functionalities/components of the intervention and comparator in the abstract. If possible, also mention theories and principles used for designing the site. Keep in mind the needs of systematic reviewers and indexers by including important synonyms. (Note: Only report in the abstract what the main paper is reporting. If this information is missing from the main body of text, consider adding it)

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Does your paper address subitem 1b-i? *

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

"For their follow-up appointments, patients received either an in-person consultation in the clinic (control group) or a video consultation with their physician (telemedicine group)."

1b-ii) Level of human involvement in the METHODS section of the ABSTRACT

Clarify the level of human involvement in the abstract, e.g., use phrases like "fully automated" vs. "therapist/nurse/care provider/physician-assisted" (mention number and expertise of providers involved, if any). (Note: Only report in the abstract what the main paper is reporting. If this information is missing from the main body of text, consider adding it)

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subitem not at all important	0	0	0	0	0	essential

Does your paper address subitem 1b-ii?

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

"For their follow-up appointments, patients received either an in-person consultation in the clinic (control group) or a video consultation with their physician (telemedicine group)."

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)

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

Does your paper address subitem 1b-iii?

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

"A prospective randomized controlled trial conducted in a German university hospital enrolled 60 patients with different knee and shoulder conditions. For their follow-up appointments, patients received either an in-person consultation in the clinic (control group) or a video consultation with their physician (telemedicine group). Patients' and physicians' subsequent evaluations of these follow-up appointments were collected and assessed using separate questionnaires."

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									
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1b-v) CONCLUSIONS/DISCUSSION in abstract for negative trials Conclusions/Discussions in abstract for negative trials: Discuss the primary outcome - if the trial is negative (primary outcome not changed), and the intervention was not used, discuss whether negative results are attributable to lack of uptake and discuss reasons. (Note: Only report in the abstract what the main paper is reporting. If this information is missing from the main body of text, consider adding it)									
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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

Does your paper address subitem 1b-v?

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

"The aim of our randomized controlled trial is to investigate whether telemedicine can be used efficiently in follow-up care for orthopedic and trauma surgery patients in Germany. To answer this question, the RCT compares an in-personal consultation in a German university hospital (Level 1 trauma center) with the use of telemedicine, namely a video consultation between physician and patient. All consultations were for follow-up care of knee and shoulder patients who displayed a variety of conditions, had previously been treated in the clinic, and were eligible to participate in the study. For their video consultation, patients did not have to travel to the clinic, but could have their follow-up appointment online regardless of their location. The subsequent evaluation of telemedicine and its efficiency focuses on patient satisfaction, physician satisfaction, and quality of care. It is hoped that studying telemedicine in broad-based use for follow-up care and analyzing its effects comprehensively will contribute to informing healthcare providers' decision-making in future."

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.

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

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

"One important tool for overcoming these challenges and guaranteeing effective healthcare in the medium to long term could be the use of telemedicine. Telemedicine offers the ability to provide medical care through real-time video consultations, without the need for personal contact and regardless of location. This could free up clinical resources, improve access to care, and increase safety for patients and medical staff. Telemedicine is already being applied successfully in various medical fields, but its use has so far been less common in orthopedic and trauma surgery. Since the outbreak of the COVID-19 pandemic, however, the need for telemedicine has risen considerably in these fields as well."

"Prior research, however, has so far left several questions unaddressed. One of them is whether the use of telemedicine is efficient not only for a restricted number of individual diseases, but also for a wider range of medical conditions. Another important question concerns the viability of deploying telemedicine under realistic conditions and cost constraints in clinical practice: It is not clear whether telemedicine remains viable for a wider range of medical conditions when offered without incurring the additional cost of human resources required to support patients in video consultations, as in an outpatient clinic. Furthermore, it is questionable whether international study results can be transferred to the German healthcare system, especially since studies show that German patients are skeptical regarding the use of telemedicine."

2b) In INTRODUCTION: Specific objectives or hypotheses

Does your paper address CONSORT subitem 2b? *

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

"The aim of our randomized controlled trial is to investigate whether telemedicine can be used efficiently in follow-up care for orthopedic and trauma surgery patients in Germany."

METHODS

3a) Description of trial design (such as parallel, factorial) including allocation ratio

Does your paper address CONSORT subitem 3a? *

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

"The study was conducted as an open, prospective, interventional, 1:1 randomized, and controlled monocenter trial at a German university hospital (University Hospital Giessen, Department of Trauma, Hand- and Reconstructive Surgery). The randomized and controlled design is based on the Consolidated Standards of Reporting Trials (CONSORT). With the parallel implementation of an intervention group, which received follow-up care through a real-time video consultation, and a control group, which received a standard follow-up consultation in the clinic, the effects of telemedicine on follow-up care were examined"

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

Does your paper address CONSORT subitem 3b? *

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

Not applicable for our study.

3b-i) Bug fixes, Downtimes, Content Change changes to methods therefore also in during the trial (e.g., major bug fixes of "unexpected events" that may have in failures/downtimes, etc. [2].	es: eheal cludes im or change	th systems portant ch s in the fu	s are ofter nanges ma nctionality	ade on the or conter	interventiont) (5-iii) ar	on or comparator nd other
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Technical irregularities in 42% of problems, sound problems, and i				onducted	l involving	g image
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

"Inclusion criteria: "(1) Patients need the ability to consent, as well as the mental and physical ability to participate in the telemedical consultation. (2) Patients' conditions require no more than a visual examination. For legal reasons, (3) a previous outpatient or inpatient stay at the clinic is required, and (4) patients have to be 18 years or older. To be able to use telemedicine, it is assumed that (5) patients own a computer, laptop, tablet or smartphone, including a microphone and camera, and (6) they have a stable internet connection. Finally, (7) patients have to speak German in order to understand the declaration of consent." As well as specified ICD-10 codes.

"Exclusion criteria: "Patients with (1) neurological diseases that do not allow the use of computer systems, and (2) patients with a diagnosis of dementia, blindness or deafness are excluded. Also, patients are excluded if they (3) have a need for in-person presence and onsite diagnostics or treatments (eg, medical imaging, laboratory, stitches, drainage) or (4) have to be touched or moved by the treating physician. This ensures that patients who require personal contact with a physician are not put at risk. Finally, (5) a lack of willingness to participate in the study or (6) the failure to consent are further exclusion criteria."

4a-i) Computer / Internet lite	eracy					
Computer / Internet literacy is often a clarified.	an implicit	"de facto"	' eligibility	criterion -	this shoul	d be explicitly
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subitem not at all important	0	0	0	0	0	essential

Does your paper address subitem 4a-i?

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

"To be able to use telemedicine, it is assumed that (5) patients own a computer, laptop, tablet or smartphone, including a microphone and camera, and (6) they have a stable internet connection."

"Patients with (1) neurological diseases that do not allow the use of computer systems, and (2) patients with a diagnosis of dementia, blindness or deafness are excluded."

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.

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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

"After initial screening for inclusion and exclusion criteria, patients were asked either at the clinic or by telephone if they would like to participate in the study during their next follow-up appointment."

4a-iii) Information giving dur	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?

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

"To be able to participate, patients had to provide informed consent after receiving written as well as oral information. Consent could be withdrawn at any time without providing reasons."

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

"At the end of the recruitment process, depending on the treatment arm, the patients received an appointment either in the clinic or for a video consultation."

4b-i) Report if outcomes wer				•	•	
Clearly report if outcomes were (self-trials) or otherwise.)assessec	d through (online que	stionnaire	s (as comr	non in web-based
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4b-ii) Report how institutional Report how institutional affiliations a affiliations with prestigious hospitals regards to an intervention. (Not a requi	re display or univers	ed to pote sities may	ntial partion	cipants [or unteer rate	es, use, and	
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Not applicable for our study.						

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5) The interventions for each group with sufficient details to allow replication,

Does your paper address sub	oitem 5-	-ii?				
Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e	uscript), o	r elaborat	e on this it	em by pro	viding add	itional
Not applicable for our study.						
5-iii) Revisions and updating						
Revisions and updating. Clearly ment (and comparator, if applicable) evaluation process, or whe Describe dynamic components such the replicability of the intervention (for	ated, or de ther the d as news fo	escribe wh levelopme eeds or ch	ether the i nt and/or o anging co	nterventio content wantent which	n underwe as "frozen"	nt major changes during the trial.
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subitem not at all important	0	0	0	0	0	essential
Does your paper address sub	oitem 5-	-iii?				
Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e	uscript), o	r elaborat	e on this it	em by pro	viding add	itional
Not applicable for our study.						
5-iv) Quality assurance meth Provide information on quality assuration provided [1], if applicable.		ods to ens	sure accura	acy and qu	uality of inf	ormation
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Does your paper address sub	oitem 5	-iv?				
Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e	uscript), c	or elaborat	e on this it	tem by pro	viding add	itional
Not applicable for our study.						
5-v) Ensure replicability by p	ublishin	g the so	urce co	de, and	or provi	ding
screenshots/screen-capture	video, a	and/or p	roviding	g flowch	arts of t	he algorithms
used		1 17	. 1.			
Ensure replicability by publishing the and/or providing flowcharts of the algorithm principle be able to replicate the students.	gorithms (used. Repl	icability (i.	e., other re		-
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Does your paper address sub						
Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e	uscript), c	or elaborat	e on this it	tem by pro	viding add	itional
Not applicable for our study.						
5-vi) Digital preservation						
Digital preservation: Provide the URL disappear over the course of the year webcitation.org, and/or publishing the pages behind login screens cannot be without login.	s; also ma e source o	ake sure th code or sci	ie interven eenshots/	ition is ard videos ald	hived (Inte	ernet Archive, e article). As
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subitem not at all important	0	0	0	0	0	essential

Does your paper address subitem 5-vi?

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.

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

O O O o essential

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

"On the day of their appointment, patients received login details for the video consultation from their physicians via text message or email. No additional registration was required for the video consultation. At the arranged time, both the attending physician and the patient logged in to the software program and the video consultation was conducted."

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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subitem not at all important O O O O essential

Does your paper address subitem 5-viii? *

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

"The online video consultation used the web-based software CLICKDOC of the German telemedicine provider CGM Mobile Services GmbH. This software is certified for and widely used in the German healthcare system. On the day of their appointment, patients received login details for the video consultation from their physicians via text message or email. No additional registration was required for the video consultation. At the arranged time, both the attending physician and the patient logged in to the software program and the video consultation was conducted. Patients were able to use a computer, laptop, tablet or smartphone to join the video consultation."

5-ix) Describe use parameters

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

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subitem not at all important	0	0	0	0	0	essential

Does your paper address subitem 5-ix?

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

"Study participants who were assigned to the intervention arm received a one-time telemedical follow-up via a real-time videoconference instead of a standard consultation in the department."

5-x) Clarify the level of human involvement

Clarify the level of human involvement (care providers or health professionals, also technical assistance) in the e-intervention or as co-intervention (detail number and expertise of professionals involved, if any, as well as "type of assistance offered, the timing and frequency of the support, how it is initiated, and the medium by which the assistance is delivered". It may be necessary to distinguish between the level of human involvement required for the trial, and the level of human involvement required for a routine application outside of a RCT setting (discuss under item 21 – generalizability).

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subitem not at all important	0	0	0	0	0	essential

Does your paper address subitem 5-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

Intervention group: "At the arranged time, both the attending physician and the patient logged in to the software program and the video consultation was conducted." Control group: "Study participants, who were assigned to the control group attended a standard follow-up consultation at the university hospital. This follow-up was conducted by the same physicians, who also treated the intervention arm."

5-xi) Report any prompts/ren Report any prompts/reminders used: use the application, what triggered th level of prompts/reminders required f application outside of a RCT setting (Clarify if tem, freque	there were ency etc. I al, and the	t may be r level of pr	necessary ompts/ren	to distingu ninders for	ish between the
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"On the day of their appointment from their physicians via text me	•		d login de	etails for	the video	consultation
5-xii) Describe any co-interv	entions	(incl. tra	ainina/sı	(troaga		
Describe any co-interventions (incl. tr addition to the targeted eHealth inter- intervention. This includes training se the level of training required for the tr RCT setting (discuss under item 21 –	aining/su vention, a essions ar ial, and th	pport): Cle s ehealth i nd support ne level of	early state intervention [1]. It may	any interv n may not v be neces	be design sary to dis	ed as stand-alone tinguish between
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information not in the ms, or briefly explain why the item is not applicable/relevant for your study

"Patients in the intervention group also received written instructions on how to perform the video consultation to minimize potential technical difficulties."

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

"The primary outcome patient satisfaction was measured using the ZAP questionnaire of the National Association of Statutory Health Insurance Physicians (NASHIP) in Germany."

"Immediately after the video consultation, patients received login details via email and were asked to evaluate the consultation via online questionnaires."

"Physician satisfaction as one of the secondary outcome parameters was assessed by questionnaires that the physicians answered following each patient consultation. The questionnaires were self-designed and differed slightly depending on the study arm."
"To be able to evaluate quality of care as a further secondary outcome, patients received the German version of the "EQ-5D-5L" questionnaire from the EuroQol Group during enrollment."
"After 3 months, the questionnaire was completed again to measure the impact of the interventions on health-related quality of life."

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

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

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	oitem 6	a-i?				
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6a-ii) Describe whether and	how "us	se" (incl	ıdina in	tensity (of use/do	osage) was
defined/measured/monitore		o (mion	aamig iii	coriorcy (31 G007G1	Joage, was
Describe whether and how "use" (inc (logins, logfile analysis, etc.). Use/ad reported in any ehealth trial.	-	-	-	•		
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6a-iii) Describe whether, hov was obtained						
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Describe whether, how, and when qua emails, feedback forms, interviews, f		ps).				
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Not applicable for our study.						
6b) Any changes to trial out	comes	after th	e trial c	ommen	ced, wit	h reasons
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Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e	uscript), c	r elaborat	e on this it	tem by pro	viding add	itional
Any changes to trial outcomes a	fter the t	rial comr	nenced, v	with reas	ons	
7a) How sample size was de NPT: When applicable, details of whe addressed			ustering b	y care pro	ovides or co	enters was
7a-i) Describe whether and h calculating the sample size	now exp	ected a	ttrition	was take	en into a	ccount when
Describe whether and how expected	attrition w	as taken i	nto accou	nt when ca	alculating t	he sample size.
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	•	_	J	7	J	
subitem not at all important	0	0	0	0	0	essential

Does your paper address subitem 7a-i?

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

"A group comparison of patient satisfaction with telemedicine in a study by Sharareh & Schwarzkopf (2014) served as the basis for our sample size calculation. This resulted in approximately 17 study participants for each group to achieve a power of 90% in a 2-sided ttest for independent samples with a global significance level of 5%. The sample size was increased by 10% for both groups to accommodate potential dropouts or withdrawals, and by another 10% to counteract a potentially skewed distribution of patient satisfaction. This resulted in a case number of 21 patients per randomization arm. To take into account the possible loss of power when using non-parametric methods, the sample size was finally increased to 30 patients per arm and thus to a total of 60 enrollments."

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

Termination criteria were established for the individual study participants. (1) The occurrence of disease symptoms that required the physical presence of the patient in the clinic. (2) Occurrence of technical problems that could not be solved in the short term. (3) Voluntary termination of the study at any time by the study participant, without giving reasons and without any disadvantage for the further treatment.

These criteria as well as unexcused absences from the consultation resulted in the recording of the study participants as dropouts/withdrawals. Patients were then treated regularly and independently of the study in the clinic.

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

"Following a 2-armed parallel group design, patients enrolled in the study were randomly assigned at a 1:1 ratio to either the intervention arm (telemedicine follow-up) or the control arm (in-person follow-up consultation in the clinic)."

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

"To ensure better balance between the arms while minimizing predictability, block randomization with randomly selected block sizes of 4, 6, and 8 was applied. One member of the study staff organized the allocation of the blocks, while a different member performed the randomization of the patients."

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

"For this purpose, sealed envelopes were used."

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

"One member of the study staff organized the allocation of the blocks, while a different member performed the randomization of the patients."

11a) If done, who was blinded after assignment to interventions (for example, participants, care providers, those assessing outcomes) and how

NPT: Whether or not administering co-interventions were blinded to group assignment

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

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

1 2 3 4 5

subitem not at all important OOOOO essential

Does your paper address subitem 11a-i? *

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

"Given that the intervention was a video consultation, blinding of physicians or patients was not possible."

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

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

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

Does your paper address subitem 11a-ii?

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

Due to the study design, patients knew whether they were in the intervention group or control group.

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

"Differences between the two study arms were analyzed using the Mann-Whitney U test or Fisher exact test, and effect sizes were reported by Pearson's correlation coefficient (r) or Cramer's V."

"In addition, the Wilcoxon signed-rank test was applied to evaluate the longitudinal data of the EQ-5D-5L VAS."

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

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

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

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

"Data were analyzed based on intention-to-treat."

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

"For patient and physician satisfaction, a subgroup analysis based on medical indication was performed."

"To examine the suitability of telemedicine for follow-up appointments in more detail, and to investigate for which patients telemedicine is most appropriate, a binary logistic regression was performed."

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

X26-i) Comment on ethics committee approval									
	1	2	3	4	5				
subitem not at all important	0	0	0	0	0	essential			

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 local ethics committee of the University of Giessen reviewed and permitted the study. In addition, the study was registered at the German Clinical Trials Register (ID:DRKS00023445)."

Outline informed concept procedures		ocedure				
Outline informed consent procedures etc.?), and what information was proviousent documents.	-				•	
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X26-iii) Safety and security p Safety and security procedures, incl. or detection of harm (e.g., education	privacy co	nsideratio			iken to red	uce the likelihood
Safety and security procedures, incl.	privacy co	nsideratio			iken to red	uce the likelihood
Safety and security procedures, incl.	privacy co and traini	onsideratio ng, availat	oility of a h	otline)		uce the likelihood essential

RESULTS

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

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

Does your paper address CONSORT subitem 13a? *

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

"A total of 60 patients agreed to participate in the study and were randomized. 30 patients were allocated to the intervention arm and 30 patients to the control arm. After randomization, 8 patients withdrew from the study. None of these patients were excluded by the physicians. Thus, 26 patients in the intervention arm and 26 patients in the control arm could be analyzed."

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

Can be found in the CONSORT flow diagram.

13b-i) Attrition diagram Strongly recommended: An attrition diagram (e.g., proportion of participants still logging in or using the intervention/comparator in each group plotted over time, similar to a survival curve) or other figures or tables demonstrating usage/dose/engagement.									
	1	2	3	4	5				
subitem not at all important	0	0	0	0	0	essential			
Does your paper address sul	oitem 13	Bb-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									
Not applicable for our study.									
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

"Patients were recruited and attended their follow-up appointments between September 2020 and April 2021. The last questionnaires for the second data collection of the EQ-5D-5L were sent out in July 2021."

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"											
	1	2	3	4	5						
subitem not at all important	0	0	0	0	0	essential					
Does your paper address subitem 14a-i? Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study While study period: COVID-19 pandemic.											
14b) Why the trial ended or	was sto	opped (e	early)								
Does your paper address CONSORT subitem 14b? * Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study Not applicable for our study.											
15) A table showing baseling group NPT: When applicable, a description centers (volume) in each group											

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									
Can be found in Table 1: Demogr	aphic ch	aracteris	tics of pa	atients.					
15-i) Report demographics a	ssociate	ed with	digital d	ivide iss	sues				
In ehealth trials it is particularly impo such as age, education, gender, socia participants, if known.		•			_				
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subitem not at all important	0	0	0	0	0	essential			
Does your paper address sul	oitem 15	5-i? *							
Copy and paste relevant sections from indicate direct quotes from your man information not in the ms, or briefly e	uscript), o	r elaborat	e on this it	tem by pro	viding add	litional			
Can be found in Table 1: Demogr	aphic ch	aracteris	tics of pa	atients.					
16) For each group, number	of part	icipants	(denor	minator`) include	ed in each			

Does your paper address CONSORT subitem 15? *

	16-i) Report multiple "denomi	nators"	and pro	ovide de	efinition	S					
	Report multiple "denominators" and provide definitions: Report N's (and effect sizes) "across a range of study participation [and use] thresholds" [1], e.g., N exposed, N consented, N used more than x times, N used more than y weeks, N participants "used" the intervention/comparator at specific pre-defined time points of interest (in absolute and relative numbers per group). Always clearly define "use" of the intervention.										
1 2 3 4 5											
	subitem not at all important	0	0	0	0	0	essential				
	Does your paper address sub	item 16	5-i? *								
	Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study										
	"In total, 100% of patients in the intervention arm and 90% (26/29) of patients in the control arm completed the questionnaires after the follow-up appointment. Of the physician questionnaires, 100% in the telemedicine group and 96% (25/26) in the control group were completed. In the intervention group, 100% of the EQ-5D-5L questionnaires were returned at baseline and 69% (18/26) after 3 months; in the control group, 88% (23/26) of the questionnaires were returned at baseline and 58% (15/26) after 3 months."										
	16-ii) Primary analysis should be intent-to-treat Primary analysis should be intent-to-treat, secondary analyses could include comparing only "users", with the appropriate caveats that this is no longer a randomized sample (see 18-i).										
		1	2	3	4	5					
	subitem not at all important	0	0	0	0	0	essential				

Does your r	paper address subitem 1	16-ii?		
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"Data were ar	nalyzed based on intention	-to-treat."		
	ch primary and second	-	_	-
Does your p	paper address CONSOR	RT subitem 17a?) *	
indicate direct	e relevant sections from the ma quotes from your manuscript), t in the ms, or briefly explain w	or elaborate on thi	s item by providing add	litional

"Although group comparison showed that patients were slightly more satisfied with telemedicine follow-up (mean 1.58) than with in-person follow-up in the clinic (mean 1.64), the difference was not statistically significant (P=.690) (Table 2)."

"The waiting time (P<.001), atmosphere (P<.001), and punctuality of the appointment (P=.002) were more satisfying for patients in the telemedicine group than in the control group, with medium to strong effects (r=.440 to r=.760)."

"A strong effect was also evident in the preference for the next follow-up appointment between the groups (V=.542). Whereas patients in the control group would prefer to visit the clinic again (64.0%), almost all patients in the telemedicine group (88.5%) would choose telemedicine for their next follow-up appointment (P<.001)."

"Physicians in the control group were significantly more satisfied with follow-up appointments (mean 1.32) than those in the telemedicine group (mean 2.42) (P=.001, r=.466), as shown in Table 3."

"However, a further group comparison, in which video consultations with technical irregularities were removed, revealed no significant group differences in physician satisfaction (mean 1.47 & 1.32, P=.310). In addition, there were no significant differences in the ability to address all relevant medical questions. For the next follow-up appointment, physicians would recommend a telemedical consultation for most patients regardless of the study arm (73.1% telemedicine group; 64.0% control group; P=.555)."

"As shown in Table 4, differences in quality of life between the groups were not significant, neither at baseline (P=.242) nor after treatment (P=.686). The difference in quality of life before and after the follow-up appointment also did not meet statistical significance between the intervention and the control arm (P=.187). In this case, the group size has changed, because only complete data sets could be considered for analysis. In both groups, the perceived average quality of life increased after treatment, although not significantly (telemedicine group: mean 69.77 to 70.44, P=.934; control group: mean 66.30 to 69.33, P=.108)."

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

Not applicable for our study.

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

Can be found in the Tables.

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

"Although group comparison showed that patients were slightly more satisfied with telemedicine follow-up (mean 1.58) than with in-person follow-up in the clinic (mean 1.64), the difference was not statistically significant (P=.690) (Table 2). This result was not affected by a subgroup analysis of the 2 medical indications, namely knee or shoulder." "Subgroup analysis showed that this difference was also significant for the treatment of shoulder patients (P=.006), but not for knee patients (P=.076). However, a further group comparison, in which video consultations with technical irregularities were removed, revealed no significant group differences in physician satisfaction (mean 1.47 & 1.32, P=.310)."

"Binary logistic regression was used to examine for which patients telemedicine follow-up is appropriate. For this purpose, the potential influence of different variables on patients' preference for their next follow-up appointment was analyzed (Table 5). The model was statistically significant $\chi 2(6) = 22.334$, P=.001, Nagelkerke`s R2 = 48.4%. It was found that neither medical indication, gender nor age had a significant influence on the choice of telemedicine. However, previous experience with video calls before the study (P=.031) and the respective study arm in which the patients were treated (P=.001) contributed significantly to predicting the choice of telemedicine."

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).

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19-i) Include privacy breached Include privacy breaches, technical pubut also incidents such as perceived unexpected/unintended incidents. "U	roblems. ⁻ or real pri	This does vacy bread	not only in ches [1], te	chnical pr	oblems, ar	nd other			
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"High number of video consultations with technical irregularities (42%, 11/26)"

Does your paper address subitem 18-i?

19-ii) Include qualitative feed staff/researchers Include qualitative feedback from pa strengths and shortcomings of the a or uses. This includes (if available) reby the developers.	rticipants pplication,	or observa	tions fron	n staff/res pint to unir	earchers, i ntended/ur	f available, on nexpected effects					
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Copy and paste relevant sections fro indicate direct quotes from your man	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 Not applicable for our study.										
DISCUSSION											
22) Interpretation consistent considering other relevant of NPT: In addition, take into account the expertise of care providers or center.	evidenc	e of the con									
22-i) Restate study questions starting with primary outcon Restate study questions and summa outcomes and process outcomes (us	nes and	process	s outcor	nes (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

"The aim of this study was to investigate whether telemedicine can be used efficiently for outpatient orthopedic and trauma surgery follow-up care in Germany at an university hospital from the perspective of patients, physicians, and the quality of care. Our data analysis shows that the use of telemedicine has no significant drawbacks compared with traditional clinical consultations in almost all aspects studied. Patients were even slightly more satisfied with telemedicine regardless of their medical condition, although the difference was not statistically significant. In addition to overall satisfaction, the authors also analyzed more specific indicators of satisfaction. Aspects such as waiting time, atmosphere and punctuality could be improved by using telemedicine."

22-ii) Highlight unanswered r	•			future r	esearch	1		
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Does your paper address sub	oitem 22	2-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								
Not applicable for our study.								

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

20-i) Typical limitations in ehealth trials

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

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

Does your paper address subitem 20-i? *

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

"The use of pen-and-paper questionnaires on the one hand and online questionnaires on the other could also have led to discrepancies. In particular, all questions had to be completed in the online questionnaires, but the same was not true for pen-and-paper questionnaires completed in the clinic. For organizational reasons, however, no uniform implementation was possible. This problem also arose for comparable studies. On the other hand, studies show that patients usually provide similar health-related answers regardless of survey formats.

Finally, when evaluating the results, it should be noted that all recruited patients consented to participating in telemedicine. This might have led to a bias in favor of higher satisfaction with telemedicine from the start, which explains why our data do not show results for the population. Short of forcing patients to participate in telemedicine, a procedure which appears both unethical in principle and unfeasible in clinical practice, there is no acceptable way of addressing this limitation."

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

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

21-i) Generalizability to othe Generalizability to other populations population, outside of a RCT setting, results for other organizations	: In particu	ılar, discus				
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"In sum, our results suggest that ideally meets several conditions assessed for each patient indiviovercoming any initial uncertain video consultation, or if problem possible, as was the case in the monitored over the long term, as satisfied with its workability."	. First, th dually, an ties. Secons os occur, a present s	e approp nd suitabl ond, if an additiona study. Th	riateness e patient y medica al in-clinic ird, techn	s of telents should all issues treatmentical irreg	nedicine s be suppo cannot be ent should gularities	should be orted in e clarified in a l always be should be
21-ii) Discuss if there were e routine application setting Discuss if there were elements in the	e RCT that	would be	different ir	n a routine	applicatio	n setting (e.g.,
prompts/reminders, more human inv impact the omission of these elemer applied outside of a RCT setting.		•				,
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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

Not applicable for our study.

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

"German Register for Clinical Trials, ID: DRKS00023445"

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

Does your paper address CONSORT subitem 24? *

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

The full trial protocol can be requested from the corresponding author.

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

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The research did not receive fu	unding fror	n an exte	ernal body	y.		
X27) Conflicts of Interest	(not a CC	ONSORT	item)			
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