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.

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Title of your manuscript * Provide the (draft) title of your manuscript.
Article Preparation Status/Stage *
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
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submitted to a journal but not reviewed yet
submitted to a journal and after receiving initial reviewer comments
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four-digit number at the end of the DOI, to be found at the bottom of each published article in JMIR)
on ms number (yet) / not (yet) submitted to / published in JMIR
Other:

TITLE AND ABSTRACT

	per address CONSORT item 1a? * ontain the phrase "Randomized Controlled Trial"? (if not, explain the reason
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Other:	
Identify the mode of in the title. Avoid a Intervention include "electronic" only if on D worlds). Use "on product names with	mode of delivery in the title of delivery. Preferably use "web-based" and/or "mobile" and/or "electronic game' mbiguous terms like "online", "virtual", "interactive". Use "Internet-based" only if es non-web-based Internet components (e.g. email), use "computer-based" or offline products are used. Use "virtual" only in the context of "virtual reality" (3- line" only in the context of "online support groups". Complement or substitute in broader terms for the class of products (such as "mobile" or "smart phone"), especially if the application runs on different platforms.
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1a-iii) Primary condition or target group in the title Mention primary condition or target group in the title, if any (e.g., "for children with Type I Diabetes") Example: A Web-based and Mobile Intervention with Telephone Support for Children with Type I Diabetes: Randomized Controlled Trial 1 2 3 4 5
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Does your paper address 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
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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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)

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2b) In INTRODUCTION: Specific objectives or hypotheses

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3b) Important changes to methods after trial commencement (such as

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4a) Eligibility criteria for participants

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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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4b) Settings and locations where the data were collected
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5) The interventions for each group with sufficient details to allow eplication, including how and when they were actually administe	
4-i) Mention names, credential, affiliations of the developers, sponsors, and owners Mention names, credential, affiliations of the developers, sponsors, and owners [6] (if uthors/evaluators are owners or developer of the software, this needs to be declared in a Conflict of interest" section or mentioned elsewhere in the manuscript).	
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5-iii) Revisions and updating Revisions and updating. Clearly mention the date and/or version number of the
application/intervention (and comparator, if applicable) evaluated, or describe whether the
intervention underwent major changes during the evaluation process, or whether the development
and/or content was "frozen" during the trial. Describe dynamic components such as news feeds o changing content which may have an impact on the replicability of the intervention (for
unexpected events see item 3b).
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5-vii) Access Access: Describe how participants accessed the application, in what setting/context, if they bay (or were paid) or not, whether they had to be a member of specific group. If known, denow 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 eviewers/readers to explore the application (also important for archiving purposes, see vi).	scribe r
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5-viii) Mode of delivery, features/functionalities/components of the intervention and	

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

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	v "use" (including intensity of use/dosage) was
lefined/measured/monitore	ed (logins, logfile analysis, etc.). Use/adoption metrics are importal
process outcomes that sho	ould be reported in any ehealth trial.
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b) Any changes easons	to trial ou	tcomes	after the tr	ial comm	enced, with
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your study
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
8a) Method used to generate the random allocation sequence
NPT: When applicable, how care providers were allocated to each trial group
3
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

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
9) Mechanism used to implement the random allocation sequence (such as sequentially numbered containers), describing any steps
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

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 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	
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 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).	
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interest" and which one will informed consent procedure				•		biases and certain expectations - discuss e.g.,
						s the "intervention of interest" and which one was
the comparator.	1	2	3	4	5	
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11b) If relevant, des	crip	otic	n c	of t	he	similarity of interventions
(this item is usually not rele intervention to a active med						s as it refers to similarity of a placebo or sham
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2a) Statistical meth econdary outcomes		use	d to	o co	ompare	groups for primary and
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2a i) Imputation tachniqu	os to d	lool v	with	ottr	ition / mis	ocina values
	al with	attriti	on /	miss	sing values	s: Not all participants will use the
						y high in ehealth trials. Specify how t from the trial were treated in the
tatistical analysis (a comple	ete cas	e ana	lysis	siss	strongly dis	scouraged, and simple imputation
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echniques such as LOCF musicipation		0	0		essential	
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your study
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
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 O O O O 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

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x26-ii) Outline informed co Outline informed consent pr Checkbox, etc.?), and what included in informed consen	ocedui inform	res e. ation	g., if was	con							be
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RESULTS
13a) For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analysed for the primary outcome
NPT: The number of care providers or centers performing the intervention in each group and the number of patients treated by each care provider in each center
Does your paper address CONSORT subitem 13a? * Copy and paste relevant sections from the manuscript (include quotes in quotation marks "like this" to indicate direct quotes from your manuscript), or elaborate on this item by providing additional information not in the ms, or briefly explain why the item is not applicable/relevant for your study
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

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additional information not i your study	n the	ms, o	r brie	elly	/ exp	nain wny	the it	em is i	пот арр	olicable/relevant for
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13b-i) Attrition diagram					,					
Strongly recommended: Ar										
the intervention/comparato		_					ne, sim	nilar to	a survi	ival curve) or other
figures or tables demonstra	ating u	ısage	/dose	e/e	enga	gement.				
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14a) Dates defining	the	per	iods	S	of r	ecruit	ment	t and	follo	w-up
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	r events"	fell ir	to th	e st	into the study period ady period, e.g., significant changes in Internet ardware or Internet delivery resources"
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ubitem not at all import	ant U			U	essential
our study					
14b) Why the trial	ended	lor	was	st	copped (early)
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aditional information no	t in the n	ns or	priet	IV P	xplain why the item is not applicable/relevant for

your study

15) A table showin for each group	g bas	eline	e de	emo	graphi	c and c	linical	characterist	ics
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15-i) Report demographi In ehealth trials it is partic issues, such as age, educ of the participants, if know	ularly im ation, ge	porta	nt to	repo	ort demog	raphics as	sociated v		
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16) For each group, number of participan each analysis and whether the analysis v	·
groups	
16-i) Report multiple "denominators" and provide define Report multiple "denominators" and provide definitions: Regarder of study participation [and use] thresholds" [1], e.g. than x times, N used more than y weeks, N participants "specific pre-defined time points of interest (in absolute an clearly define "use" of the intervention.	eport N's (and effect sizes) "across a , N exposed, N consented, N used more used" the intervention/comparator at
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Does your paper address subitem 16-i? * Copy and paste relevant sections from the manuscript (in this" to indicate direct quotes from your manuscript), or el additional information not in the ms, or briefly explain why your study	aborate on this item by providing
16-ii) Primary analysis should be intent-to-treat Primary analysis should be intent-to-treat, secondary ana	
"users", with the appropriate caveats that this is no longe 1 2 3 4 5	r a randomized sample (see 18-i).
subitem not at all important 0 0 0 0 essentia	-
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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
17a-i) Presentation of process outcomes such as metrics of use and intensity of use In addition to primary/secondary (clinical) outcomes, the presentation of process outcomes such as metrics of use and intensity of use (dose, exposure) and their operational definitions is critical. This does not only refer to metrics of attrition (13-b) (often a binary variable), but also to more continuous exposure metrics such as "average session length". These must be accompanied by a technical description how a metric like a "session" is defined (e.g., timeout after idle time) [1] (report under item 6a).
1 2 3 4 5
subitem not at all important O O O O 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	
17b) For binary outcomes, presentation of both absolute and relative effect sizes is recommended	<u>}</u>
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	
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	

	pari	ng o	nly	user	s is	users not uncommon in ehealth trials, but if done, it mple and no longer an unbiased sample from a
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19) All important ha	rms	s o	r u	nin	ten	nded effects in each group
(for specific guidance see C	ONS	SOR	T fo	r ha	rms	3)

Does your paper address CONSORT subitem 19? *

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

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19-i) Include privacy brea Include privacy breaches, to participants, but also incide and other unexpected/unint effects [2].	echnica nts suc	l prob h as	olem perc	s. T	his does ned or real p	orivacy brea	aches [1], 1	technical pr	roblems,
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19-ii) Include qualitative f Include qualitative feedback									
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DISCUSSION										
22) Interpretation coharms, and conside						-		_	bene	fits and
NPT: In addition, take into a unequal expertise of care p								lack of	f or part	ial blinding, and
22-i) Restate study questi- with primary outcomes an Restate study questions and outcomes and process outcomes	nd pr d sur	roces mma	ss o	utc	ome	s (use)				
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22-ii) Highlight unanswered new questions, suggest future research Highlight unanswered new questions, suggest future research.

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nd, if relevant, multip 0-i) Typical limitations in elypical limitations in ehealth to	pli hea	cit alth	y o	of a	na nts	ources of potential bias, imprecision lyses in ehealth trials are rarely blinded. Ehealth trial and risk for a Type I error. Discuss biases due to
						es through informed consent procedures,
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21) Generalisability (external validity, applicability) of the trial findings

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

21-i) Generalizability to other populations

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

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Does your paper address subitem 21-i?

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

//

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

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

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subitem not at all important	0	0	0	0	0	essential

Does your paper address subitem 21-ii?

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

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
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

oes your paper address CONSORT subitem 25? * opy and paste relevant sections from the manuscript (include quotes in quotation marks "lidis" 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 our study	
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	ıg
5) Sources of funding and other support (such as supply of drug ble of funders	ugs)

Does your paper address subitem X27-i?

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

your study
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