

Online Supplement. Preferred Reporting Items for Systematic Review and Meta-Analysis Protocols (PRISMA-P) flow diagram of our search history

Table S1. Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) Systematic Review Checklist

Checklist			
Selection/Topic	#	Checklist Item	Reported on Page #
TITLE			
Title	1	Identify the report as a systematic review, meta-analysis, or	1
		both.	
ABSTRACT			
Structured summary	2	Provide a structured summary including, as applicable:	2
		background; objectives; data sources; study eligibility	
		criteria, participants, and interventions; study appraisal and	
		synthesis methods; results; limitations; conclusions	
		and implications of key findings; systematic review	
		registration number.	
INTRODUCTION			
Rationale	3	Describe the rationale for the review in the context of what is	5
		already known.	
Objectives	4	Provide an explicit statement of questions being addressed	5
		with reference to participants, interventions,	
		comparisons, outcomes, and study design (PICOS).	
METHODS			
Protocol and	5	Indicate if a review protocol exists, if and where it can be	N/A
registration		accessed (e.g., Web address), and, if available, provide	
		registration information including registration number.	
Eligibility criteria	6	Specify study characteristics (e.g., PICOS, length of follow-	6
		up) and report characteristics (e.g., years considered,	
		language, publication status) used as criteria for eligibility,	
		giving rationale.	
Information sources	7	Describe all information sources (e.g., databases with dates	5
		of coverage, contact with study authors to identify	
		additional studies) in the search and date last searched.	
Search	8	Present full electronic search strategy for at least one	Online Supplement
		database, including any limits used, such that it could be	
		repeated.	
Study selection	9	State the process for selecting studies (i.e., screening,	6
		eligibility, included in systematic review, and, if applicable,	
		included in the meta-analysis).	
Data collection	10	Describe method of data extraction from reports (e.g.,	6
process		piloted forms, independently, in duplicate) and any	
		processes for obtaining and confirming data from	
		investigators.	
Data items	11	List and define all variables for which data were sought (e.g.,	6
		PICOS, funding sources) and any assumptions and	
		simplifications made.	
Risk of bias in	12	Describe methods used for assessing risk of bias of	7
individual studies		individual studies (including specification of whether this	
		was done at the study or outcome level), and how this	
		information is to be used in any data synthesis.	
Summary measures	13	State the principal summary measures (e.g., risk ratio,	N/A
,		1 1	

		difference in means).	
Synthesis of results	14	Describe the methods of handling data and combining results	N/A
·		of studies, if done, including measures of	
		consistency (e.g., I2) for each meta-analysis.	
Risk of bias across	15	Specify any assessment of risk of bias that may affect the	N/A
studies		cumulative evidence (e.g., publication bias, selective	
		reporting within studies).	
Additional analyses	16	Describe methods of additional analyses (e.g., sensitivity or	N/A
,		subgroup analyses, meta-regression), if done,	
		indicating which were pre-specified.	
RESULTS			
Study selection	17	Give numbers of studies screened, assessed for eligibility,	7
		and included in the review, with reasons for exclusions	
		at each stage, ideally with a flow diagram.	
Study characteristics	18	For each study, present characteristics for which data were	7-20
		extracted (e.g., study size, PICOS, follow-up period)	
		and provide the citations.	
Risk of bias within	19	Present data on risk of bias of each study and, if available,	7-20
studies		any outcome-level assessment (see Item 12).	
Results of individual	20	For all outcomes considered (benefits or harms), present, for	N/A
studies		each study: (a) simple summary data for each	
		intervention group and (b) effect estimates and confidence	
		intervals, ideally with a forest plot.	
Synthesis of results	21	Present results of each meta-analysis done, including	N/A
		confidence intervals and measures of consistency.	
Risk of bias across	22	Present results of any assessment of risk of bias across	N/A
studies		studies (see Item 15).	
Additional analysis	23	Give results of additional analyses, if done (e.g., sensitivity	N/A
		or subgroup analyses, meta-regression [see Item 16]).	
DISCUSSION			
Summary of	24	Summarize the main findings including the strength of	22
evidence		evidence for each main outcome; consider their	
		relevance to key groups (e.g., health care providers, users,	
		and policy makers).	
Limitations	25	Discuss limitations at study and outcome level (e.g., risk of	25-27
		bias), and at review level (e.g., incomplete retrieval of	
		identified research, reporting bias).	
Conclusions	26	Provide a general interpretation of the results in the context	27
		of other evidence, and implications for future	
		research.	
FUNDING			
Funding	27	Describe sources of funding for the systematic review and	N/A
		other support (e.g., supply of data); role of funders for	
		the systematic review.	

## Supplement 1

## Sample Search Strategy: PubMed

- 01. Internet
- 02. Internet-based
- 03. web-based
- 04. mobile app\*
- 05. smartphone app\*
- 06. technology
- 07. therapist OR clinician
- 08. training OR education
- 09. 1 or 2 or 3 or 4 or 5 or 6
- 11. 07 and 08.
- 12. limit to (English language)
- 13. limit to (peer reviewed)