

### 3. Tabellen, Extraktionsbögen und weitere Unterlagen zu den Anträgen

#### 3.3 Studienextraktionsbogen für systematische Übersichten

Nr.	Feld	Hinweise
1.	<b>Autor</b>	
2.	<b>Titel</b>	
3.	<b>Quelle</b>	
4.	<b>Bezugsrahmen</b>	
5.	<b>Fragestellung/Zielsetzung</b>	
6.	<b>Krankheit</b>	
7.	<b>Intervention (Angaben zur Zusammensetzung nach Art und Menge)</b>	
8.	<b>Einschlusskriterien</b>	The selection criteria (i. e. population, intervention, outcome, and study design): methods for validity assessment, data abstraction, and study characteristics, and quantitative data synthesis in sufficient detail to permit replication
9.	<b>Ausschlusskriterien</b>	
10.	<b>Ergebnis der Recherche</b>	Characteristics of the RCTs included and excluded: qualitative and quantitative findings (i. e. point estimates and confidence intervals); and sub-group analyses
11.	<b>Einführung</b>	The explicit clinical problem, biological rationale for the intervention and rationale for the review
12.	<b>Beschreibung der Suche</b>	The information sources in detail (e. g. databases, registers, personal files, expert informants, agencies, hand-searching), and any restrictions (years considered, publication status, language of publication)
13.	<b>Methodische Beschreibung des Vorgehens</b>	Data abstraction:  The process or processes used (e. g. completed independently, in duplicate)

<b>14.</b>	<b>Validität</b>	The criteria and process used (e. g. masked conditions, quality assessment, and their findings)
<b>15.</b>	<b>Charakterisierung der Studien</b>	The type of study design, participants' characteristics, details of intervention, outcome definitions, and how clinical heterogeneity was assessed
<b>16.</b>	<b>Quantitative Ergebnisse der Synthese</b>	The principal measures of effect (e.g. relative risk), method of combining results (statistical testing and confidence intervals), handling of missing data; how statistical heterogeneity was assessed; a rationale for any priori sensitivity and sub-group analyses; and any assessment of publication bias
<b>17.</b>	<b>Darstellung in einem Flussdiagramm</b>	Trial flow: Provide a meta-analysis profile summarising trial flow
<b>18.</b>	<b>Charakterisierung der gefundenen Studien</b>	Present descriptive data for each trial (e.g. age, sample size, intervention, dose, duration, follow-up period)
<b>19.</b>	<b>Nebenwirkungen</b>	
<b>20.</b>	<b>Schlussfolgerung</b>	The main results
<b>21.</b>	<b>Bewertung der methodischen Qualität</b>	Summarise key findings; discuss clinical inferences based on internal and external validity; interpret the results in the light of the totality of available evidence; describe potential biases in the review process e. g. publication bias); and suggest a future re-search agenda