

Instructions for evaluating the reliability and relevance of *in vitro* toxicity studies using the SciRAP tool.

Note: this guidance also applies to use of the SciRAP in vitro tool for evaluating in vitro studies on nano materials.

Introduction:

The SciRAP tools for evaluating *in vitro* studies and *in vitro* studies on nano materials allow for evaluation of reliability (divided into reporting quality and methodological quality) and relevance. The evaluation often has to be endpoint-specific, meaning that the evaluation is carried out focusing on one of several endpoints/effects investigated in the study. Separate evaluations may thus be necessary for different endpoints in one study.

Download the excel file containing the assessment sheet available on the SciRAP website. The assessment sheet contains pre-defined criteria/items to be evaluated in 3 sections for **reporting quality**, **methodological quality**, and **relevance**. The reporting and methodological quality sections are divided in specific categories: Test compound and controls, Test system, Administration of the test compound, Data collection and analysis, Funding and competing interests (only in the reporting quality section), and Other (**Fig. 1**). The SciRAP tool for studies on nano materials also contains a category of reporting quality criteria for Physicochemical properties of the test item.

	METHODOLOGICAL QUALITY	SELECTION	COMMENT
Test compound and controls			
1	The test compound or mixture was unlikely to contain any impurities that may significantly have affected the results of the study.		
2	It was likely that the test compound was soluble at the concentrations used.		
3	An appropriate solvent (vehicle) was used that is not expected to interfere with the results of the study at the concentration used.		
4	A solvent (vehicle) control was included.		
5	An appropriate positive control was included, and the expected result was observed from this treatment.		
Test System			
6	A reliable and sensitive test system (e.g., cell line / cells/ tissue / organ / embryo / sub-cellular fractions) with metabolic competence, if relevant, was used for investigating the test compound and endpoints.		
7	Conditions for cultivation and/or maintenance of the cell line / cells / tissue / organ / embryo / sub-cellular fractions (incubation temperature, humidity, CO2 concentration, media used, number of cell passages, control of contamination) were appropriate.		
Administration of the test compound			
8	The duration of exposure was suitable for the test system and investigated endpoints.		
9	The concentrations used were suitable for the test system and investigated endpoints.		
10	The test conditions during and after exposure to the test compound were suitable (media and serum used, cell density, incubation temperature, humidity, CO2 concentration).		
Data collection and analysis			
11	Reliable and sensitive tests and/or analytical methods were used for investigating the endpoints.		
12	Sufficient numbers of replicates or repetitions of the experiment were used to generate reliable and valid results.		
13	Measurements were collected at suitable time points in order to generate sensitive, valid and reliable data.		
14	Cytotoxicity was measured and the test compound did not cause cytotoxicity that significantly affected the results.		
15	The statistical methods were clearly described and do not seem inappropriate, unusual or unfamiliar.		
Other			
16	Are there any other aspects of study design, performance or reporting that influence reliability?		

Fig. 1 Categories of criteria in Methodological Quality section of the SciRAP tool.

Evaluation may be conducted for either reporting quality, methodological quality, or relevance, or all three, depending on the purpose of evaluation. Although not required, evaluating reporting quality of the study before moving into the evaluation of methodological quality and relevance may in some cases save time and resources as it allows for identification of studies that have obvious deficiencies in reporting, hampering further evaluation.

Evaluation of the criteria:

When you evaluate the criteria/items, choose one of the options from the drop-down menu in the "SELECTION" column (fulfilled, partially fulfilled, or not fulfilled for reporting and methodological quality, directly relevant, indirectly relevant, or not relevant in the relevance section, (Fig. 2). This drop-down menu is in almost every cell in the "SELECTION" column.

	METHODOLOGICAL QUALITY	SELECTION	COMMENT
Test compound and controls			
1	The test compound or mixture was unlikely to contain any impurities that may significantly have affected the results of the study.	fulfilled	
2	It was likely that the test compound was soluble at the concentrations used.	partially fulfilled	
3	An appropriate solvent (vehicle) was used that is not expected to interfere with the results of the study at the concentration used.	not fulfilled	
4	A solvent (vehicle) control was included.	not reported	
5	An appropriate positive control was included, and the expected result was observed from this treatment.	REMOVE	
Test System			
6	A reliable and sensitive test system (e.g., cell line / cells/ tissue / organ / embryo / sub-cellular fractions) with metabolic competence, if relevant, was used for investigating the test compound and endpoints.	fulfilled	
7	Conditions for cultivation and/or maintenance of the cell line / cells / tissue / organ / embryo / sub-cellular fractions (incubation temperature, humidity, CO2 concentration, media used, number of cell passages, control of contamination) were appropriate.	partially fulfilled	
Administration of the test compound			
8	The duration of exposure was suitable for the test system and investigated endpoints.	not reported	
9	The concentrations used were suitable for the test system and investigated endpoints.	not reported	

Fig. 2 Drop-down menu for the criteria in Methodological Quality section of the SciRAP tool.

Guidance for evaluating individual methodological quality criteria and relevance items is available by pointing to the criterion with the cursor (the criterion containing the guidance has a red right corner, Fig. 3).

	METHODOLOGICAL QUALITY	SELECTION	COMMENT
Test compound and controls			
1	The test compound or mixture was unlikely to contain any impurities that may significantly have affected the results of the study.	fulfilled	Guidance: Purity of the test compound, or the composition of substances in a mixture can potentially affect study results. Purity and composition is also an important aspect to consider in terms of the relevance of the test compound to the compound being risk assessed. Ideally, in the case of single compounds, the test chemical should be of the highest available purity. Significant impurities, or isomers of the test compound, are more likely to be present, and/or to impact toxicity for certain compounds. For example, PCBs (individual or in mixtures) are often contaminated with low levels of potentially highly toxic dioxins. The measured toxicity of the test compound may then be due to the contaminant. In such cases information about the level of purity and composition is critical. How to judge this criterion: Fulfilled – The test compound has been clearly identified and characterized and is of sufficient purity. In cases of mixtures, the composition of substances is well characterized and their individual purities are sufficient. Partially fulfilled – The purity of the test compound has not been described but it is unlikely that impurities are present that would significantly affect the results of the study. Not fulfilled – The test compound or mixture is likely to contain impurities that can affect study results.
2	It was likely that the test compound was soluble at the concentrations used.	partially fulfilled	
3	An appropriate solvent (vehicle) was used that is not expected to interfere with the results of the study at the concentration used.	not fulfilled	
4	A solvent (vehicle) control was included.	not reported	
5	An appropriate positive control was included, and the expected result was observed from this treatment.	REMOVE	
Test System			
6	A reliable and sensitive test system (e.g., cell line / cells/ tissue / organ / embryo / sub-cellular fractions) with metabolic competence, if relevant, was used for investigating the test compound and endpoints.	fulfilled	
7	Conditions for cultivation and/or maintenance of the cell line / cells / tissue / organ / embryo / sub-cellular fractions (incubation temperature, humidity, CO2 concentration, media used, number of cell passages, control of contamination) were appropriate.	partially fulfilled	
Administration of the test compound			
8	The duration of exposure was suitable for the test system and investigated endpoints.	not fulfilled	
9	The concentrations used were suitable for the test system and investigated endpoints.	not reported	
10	The test conditions during and after exposure to the test compound were suitable (media and serum used, cell density, incubation temperature, humidity, CO2 concentration).	REMOVE	
Data collection and analysis			

Fig. 3 Guidance for evaluating criteria in the SciRAP tool.

Criterion no. 24 in the reporting quality section and criterion no. 16 in methodological quality section of the SciRAP *in vitro* tool, and no. 38 and 19 in the tool for studies on nano materials, provide space for free text comments on additional aspects that affect study reliability. These criteria do not contain the drop-down menu with options.

You may use the "COMMENT" column to write free text comments, for example explaining your evaluation of a specific criterion (Fig. 4).

	METHODOLOGICAL QUALITY	SELECTION	COMMENT
Test compound and controls			
1	The test compound or mixture was unlikely to contain any impurities that may significantly have affected the results of the study.	fulfilled	
2	It was likely that the test compound was soluble at the concentrations used.	partially fulfilled	
3	An appropriate solvent (vehicle) was used that is not expected to interfere with the results of the study at the concentration used.	not fulfilled	
4	A solvent (vehicle) control was included.	not reported	WRITE A NOTE!
5	An appropriate positive control was included, and the expected result was observed from this treatment.	REMOVE	
Test System			
6	A reliable and sensitive test system (e.g., cell line / cells/ tissue / organ / embryo / sub-cellular fractions) with metabolic competence, if relevant, was used for investigating the test compound and endpoints.	fulfilled	

Fig. 4 Writing a note in the "COMMENT" column.

Judging criteria as “not reported”

If a criterion cannot be judged, you can select the option “not reported” in the drop-down menu (**Fig. 2**). This is primarily intended for methodological quality criteria when sufficient information is lacking to make a judgment regarding whether the criterion is fulfilled or not. Note that for reporting quality, if information is missing you should select “not fulfilled”.

Removing criteria

Individual criteria may be considered more or less critical in the specific case you are working on, and the SciRAP tool includes a function to remove criteria for reporting and methodological quality. In that case, choose "REMOVE" in the drop-down menu of the "SELECTION" column instead of fulfilled, partially fulfilled, not fulfilled (**Fig. 2**). Removed criteria will not be included in the colour profile or % fulfilled criteria calculation. Motivations for removing criteria can be provided in the "COMMENT" column (**Fig. 4**).

NOTE: removing criteria will have an impact on the colour profile and the % fulfilled criteria. It is therefore important that the same criteria are removed in evaluations that are going to be compared to each other. Items in the Relevance section cannot be removed.

Interpreting the results of the SciRAP tool:

Results of the study assessment are shown right below the relevance section of the SciRAP tool in the form of % fulfilled criteria, as well as a colour profile.

	% FULFILLED CRITERIA	
	REPORTING	METHOD
Study overall	50.00	37.50
Test compounds and controls	62.50	37.50
Test system	40.00	75.00
Administration of test compound	50.00	0.00
Data collection and analysis	62.50	37.50
Funding and competing interests	25.00	

Fig. 5 Table with % fulfilled criteria.

Percent fulfilled criteria

The results show % fulfilled criteria of for the study overall, as well as for the specific criteria categories (**Fig. 5**).

- The % fulfilled criteria is calculated as follows:

$$\text{SciRAP score (\%)} = \frac{F+(PF*0.5)}{T} * 100\% / \text{SciRAP score (\%)} = \frac{DR+(IR*0.5)}{T} * 100\%$$

where *F* is the number of fulfilled criteria, *PF* is the number of partially fulfilled criteria, and *T* is the total number of criteria. In other words, partially fulfilled criteria contribute half the value as fulfilled criteria. Criteria that have been removed are excluded from the calculation.

The % fulfilled criteria can have a value ranging from 0 (all criteria are judged as "not fulfilled"/"not reported") to 100 (all criteria are judged as "fulfilled").

NOTE:

- selecting “not reported” for a criterion will have the same impact as “not fulfilled” on the % fulfilled value. The user should take care to note the reason for leaving a criterion as "not reported".
- removing criteria will have an impact on the % fulfilled criteria, as well as the colour profile. It is therefore important that the same criteria are removed in evaluations that are going to be compared to each other.
- importantly, the % fulfilled criteria cannot be considered on its own but should be interpreted together with the colour profile when concluding on study reliability. The colour profile is crucial to identify where a study's strengths and weaknesses lie and is more informative than the % fulfilled criteria for this purpose.

Colour profile

In the colour profile, the evaluations of reliability and relevance are illustrated in bar charts (**Fig. 6**), showing green for fulfilled criteria, yellow for partially fulfilled and red for criteria that were not fulfilled. Criteria that were "not reported" will be shown as grey. Relevance items evaluated as relevant are shown as green, indirectly relevant items are shown as yellow, and if the item was evaluated as being not relevant for the risk assessment or problem formulation, it is shown as red. The bar charts do not include criteria that have been removed.

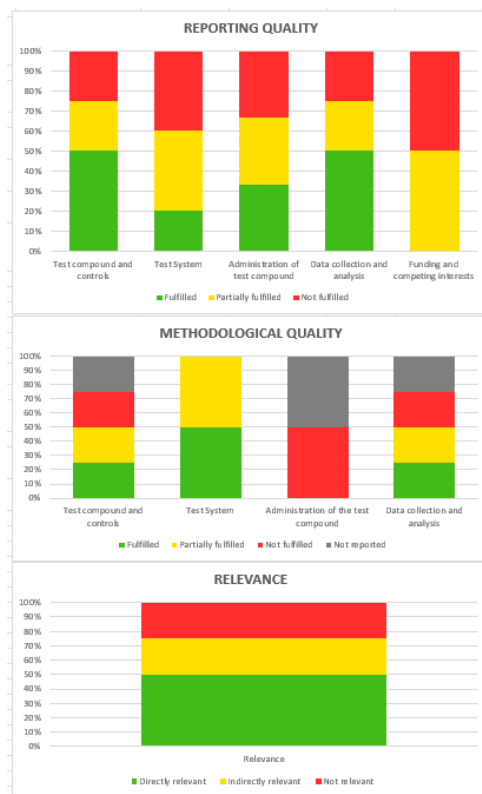


Fig. 6 The evaluations of reliability and relevance are illustrated in bar charts.

Categorisation of reliability and relevance

The SciRAP tool does not provide cut-off values or a pre-defined scheme for categorisation of the reliability and relevance of *in vivo* toxicity data. Principles for such categorisation needs to be established on a case-by-case basis and should be fit for purpose for the assessment at hand. Some examples of how the output of the SciRAP evaluation can be used in different contexts, including weight of evidence assessment, are provided in published articles. For example:

Holmer ML, Zilliacus J, Draskau MK, Hlisníková H, Beronius A, Svingen T. 2024. Methodology for developing data-rich Key Event Relationships for Adverse Outcome Pathways exemplified by linking decreased androgen receptor activity with decreased anogenital distance. *Reprod Toxicol.* 128:108662. doi: 10.1016/j.reprotox.2024.108662. Epub ahead of print. PMID: 38986849.

Röhl C, Batke M, Damm G, Freyberger A, Gebel T, Gundert-Remy U, Hengstler JG, Mangerich A, Matthiessen A, Partosch F, Schupp T, Wollin KM, Foth H. 2022. New aspects in deriving health-based guidance values for bromate in swimming pool water. *Arch Toxicol.* 96(6):1623-1659. doi: 10.1007/s00204-022-03255-9. PMID: 35386057; PMCID: PMC9095538.

Wiklund L and Beronius A. 2022. Systematic evaluation of the evidence for identification of endocrine disrupting properties of Bisphenol F. *Toxicology.* 476:153255. doi: <https://doi.org/10.1016/j.tox.2022.153255>

If you have any questions, please do not hesitate to contact us at anna.beronius@ki.se.