PRACTICAL APPLICATION



Preferred Reporting Items for Studies Mapping onto Preference-Based Outcome Measures: The MAPS Statement

Stavros Petrou¹ · Oliver Rivero-Arias² · Helen Dakin³ · Louise Longworth⁴ · Mark Oppe⁵ · Robert Froud^{1,6} · Alastair Gray³

Published online: 1 August 2015

© The Author(s) 2015. This article is published with open access at Springerlink.com

Abstract 'Mapping' onto generic preference-based outcome measures is increasingly being used as a means of generating health utilities for use within health economic evaluations. Despite the publication of technical guides for the conduct of mapping research, guidance for the reporting of mapping studies is currently lacking. The MAPS (MApping onto Preference-based measures reporting Standards) statement is a new checklist, which aims to promote complete and transparent reporting of mapping studies. The primary audiences for the MAPS statement are researchers reporting mapping studies, the funders of the research, and peer reviewers and editors involved in assessing mapping studies for publication. A de novo list of 29 candidate reporting items and accompanying explanations was created by a working group comprising six health economists and one Delphi methodologist. Following a two-round modified Delphi survey with representatives from academia, consultancy, health technology assessment agencies and the biomedical journal editorial community, a final set of 23 items deemed essential for transparent reporting, and accompanying explanations, was developed.

The items are contained in a user-friendly 23-item check-list. They are presented numerically and categorised within six sections, namely: (1) title and abstract; (2) introduction; (3) methods; (4) results; (5) discussion; and (6) other. The MAPS statement is best applied in conjunction with the accompanying MAPS explanation and elaboration document. It is anticipated that the MAPS statement will improve the clarity, transparency and completeness of reporting of mapping studies. To facilitate dissemination and uptake, the MAPS statement is being co-published by seven health economics and quality-of-life journals, and broader endorsement is encouraged. The MAPS working group plans to assess the need for an update of the reporting checklist in 5 years' time.

Key Points

This paper summarises the development of the MAPS reporting statement, a checklist of essential items that authors should consider when reporting mapping studies.

It is anticipated that the MAPS reporting statement will promote clarity, transparency and completeness of reporting of mapping studies.

- Warwick Clinical Trials Unit, Warwick Medical School, University of Warwick, Coventry CV4 7AL, UK
- National Perinatal Epidemiology Unit, Nuffield Department of Population Health, University of Oxford, Oxford, UK
- Nuffield Department of Population Health, Health Economics Research Centre, University of Oxford, Oxford, UK
- ⁴ Health Economics Research Group, Brunel University London, Uxbridge, UK
- ⁵ EuroQol Research Foundation, Rotterdam, The Netherlands
- Norges Helsehøyskole, Campus Kristiania, Oslo, Norway

1 Introduction

The process of 'mapping' onto generic preference-based outcome measures is increasingly being used as a means of generating health utilities for application within health economic evaluations [1]. Mapping involves the

986 S. Petrou et al.

development and use of an algorithm (or algorithms) to predict the primary outputs of generic preference-based outcome measures, i.e. health utility values, using data on other indicators or measures of health. The source predictive measure may be a non-preference based indicator or measure of health outcome or, more exceptionally, a preference-based outcome measure that is not preferred by the local health technology assessment agency. The algorithm(s) can subsequently be applied to data from clinical trials, observational studies or economic models containing the source predictive measure(s) to predict health utility values in contexts where the target generic preferencebased measure is absent. The predicted health utility values can then be analysed using standard methods for individual-level data (e.g. within a trial-based economic evaluation), or summarised for each health state within a decision-analytic model.

Over recent years there has been a rapid increase in the publication of studies that use mapping techniques to predict health utility values, and databases of published studies in this field are beginning to emerge [2]. Some authors [3] and agencies [4] concerned with technology appraisals have issued technical guides for the conduct of mapping research. However, guidance for the reporting of mapping studies is currently lacking. In keeping with health-related research more broadly [5], mapping studies should be reported fully and transparently to allow readers to assess the relative merits of the investigation [6]. Moreover, there may be significant opportunity costs associated with regulatory and reimbursement decisions for new technologies informed by misleading findings from mapping studies. This has led to the development of the MAPS (MApping onto Preference-based measures reporting Standards) statement, which we summarise in this paper.

The aim of the MAPS statement is to provide recommendations, in the form of a checklist of essential items, which authors should consider when reporting a mapping study. It is anticipated that the checklist will promote complete and transparent reporting by researchers. The focus, therefore, is on promoting the quality of reporting of mapping studies, rather than the quality of their conduct, although it is possible that the reporting statement will also indirectly enhance the methodological rigour of the research [7]. The MAPS statement is primarily targeted at researchers developing mapping algorithms, the funders of the research, and peer reviewers and editors involved in the manuscript review process for mapping studies [5, 6]. In developing the reporting statement, the term 'mapping' is used to cover all approaches that predict the outputs of generic preference-based outcome measures using data on other indicators or measures of health, and encompasses related forms of nomenclature used by some researchers, such as 'cross-walking' or 'transfer to utility' [1, 8].

Similarly, the term 'algorithm' is used in its broadest sense to encompass statistical associations and more complex series of operations.

2 The Development of the MAPS Statement

The development of the MAPS statement was informed by recently published guidance for health research reporting guidelines [5] and broadly modelled other recent reporting guideline developments [9–14]. A working group comprising six health economists (SP, ORA, HD, LL, MO, AG) and one Delphi methodologist (RF) was formed following a request from an academic journal to develop a reporting statement for mapping studies. One of the working group members (HD) had previously conducted a systematic review of studies mapping from clinical or health-related quality-of-life measures onto the EQ-5D [2]. Using the search terms from this systematic review, as well as other relevant articles and reports already in our possession, a broad search for reporting guidelines for mapping studies was conducted. This confirmed that no previous reporting guidance had been published. The working group members therefore developed a preliminary de novo list of 29 reporting items and accompanying explanations. Following further review by the working group members, this was subsequently distilled into a list of 25 reporting items and accompanying explanations.

Members of the working group identified 62 possible candidates for a Delphi panel from a pool of active researchers and stakeholders in this field. The candidates included individuals from academic and consultancy settings with considerable experience in mapping research, representatives from health technology assessment agencies that routinely appraise evidence informed by mapping studies, and biomedical journal editors. Health economists from the MAPS working group were included in the Delphi panel. A total of 48 of the 62 (77.4 %) individuals agreed to participate in a Delphi survey aimed at developing a minimum set of standard reporting requirements for mapping studies with an accompanying reporting checklist.

The Delphi panellists were sent a personalised link to a Web-based survey, which had been piloted by members of the working group. Non-responders were sent up to two reminders after 14 and 21 days. The panellists were anonymous to each other throughout the study and their identities were known only to one member of the working group. The panellists were invited to rate the importance of each of the 25 candidate reporting items identified by the working group on a 9-point rating scale (1, "not important", to 9, "extremely important"); describe their confidence in their ratings ("not confident", "somewhat confident" or "very confident"); comment on the candidate

The MAPS Statement 987

items and their explanations; suggest additional items for consideration by the panellists in subsequent rounds; and to provide any other general comments. The candidate reporting items were ordered within six sections: (1) title and abstract; (2) introduction; (3) methods; (4) results; (5) discussion; and (6) other. The panellists also provided information about their geographical area of work, sex, and primary and additional work environments. The data were imported into Stata (version 13; Stata-Corp, College Station, TX, USA) for analysis.

A modified version of the Research ANd Development (RAND)/University of California Los Angeles (UCLA) appropriateness method was used to analyse the round one responses [15]. This involved calculating the median score, the inter-percentile range (IPR) (30th and 70th), and the inter-percentile range adjusted for symmetry (IPRAS), for each item (i) being rated. The IPRAS includes a correction factor for asymmetric ratings, and panel disagreement was judged to be present in cases if $IPR_i > IPRAS_i$ [15]. We modified the RAND/UCLA approach by asking panellists about 'importance' rather than 'appropriateness' per se. Assessment of importance followed the classic RAND/ UCLA definitions, categorised simply as whether the median rating fell between 1 and 3 (unimportant), 4 and 6 (neither unimportant nor important), or 7 and 9 (important) [15].

The results of round one of the Delphi survey were reviewed at a face-to-face meeting of the working group. A total of 46 of the 48 (95.8 %) individuals who agreed to participate completed round one of the survey. Of the 25 items, 24 were rated as important, with one item ("Source of Funding") rated as neither unimportant nor important. There was no evidence of disagreement on ratings of any items according to the RAND/UCLA method. These findings did not change when the responses of the MAPS working group were excluded. Based on the qualitative feedback received in round one, items describing "Modelling Approaches" and "Repeated Measurements" were merged, as were items describing "Model Diagnostics" and "Model Plausibility". In addition, amendments to the wording of several recommendations and their explanations were made in the light of qualitative feedback from the panellists.

Panellists participating in round one were invited to participate in a second round of the Delphi survey. A summary of revisions made following round one was provided. This included a document in which revisions to each of the recommendations and explanations were displayed in the form of track changes. Panellists participating in round two were provided with group outputs (mean scores and their standard deviations, median scores and their IPRs, histograms and RAND/UCLA labels of importance and agreement level) summarising the round

one results (and disaggregated outputs for the merged items). They were also able to view their own round one scores for each item (and disaggregated scores for the merged items). Panellists participating in round two were offered the opportunity to revise their rating of the importance of each of the items and informed that their rating from round one would otherwise hold. For the merged items, new ratings were solicited. Panellists participating in round two were also offered the opportunity to provide any further comments on each item or any further information that might be helpful to the group. Non-responders to the second round of the Delphi survey were sent up to two reminders after 14 and 21 days. The analytical methods for the round two data mirrored those for the first round.

The results of the second round of the Delphi survey were reviewed at a face-to-face meeting of the working group. A total of 39 of the 46 (84.8 %) panellists participating in round one completed round two of the survey. All 23 items included in the second round were rated as important with no evidence of disagreement on ratings of any items according to the RAND/UCLA method. Qualitative feedback from the panellists participating in round two led to minor modifications to wording of a small number of recommendations and their explanations. This was fed back to the round two respondents who were given a final opportunity to comment on the readability of the final set of recommendations and explanations. Based on these methods, a final consensus list of 23 reporting items was developed.

3 The MAPS Statement

The MAPS statement is a 23-item checklist of recommendations (Table 1) that we consider essential for complete and transparent reporting of studies that map onto generic preference-based outcome measures. The 23 reporting items are presented numerically and categorised within six sections, namely: (1) title and abstract (two items); (2) introduction (two items); (3) methods (nine items); (4) results (six items); (5) discussion (three items); and (6) other (one item). The reporting of each item does not necessarily have to follow the order within the MAPS statement. Rather, what is important is that each recommendation is addressed either in the main body of the report or its appendices. Several biomedical journals have endorsed the MAPS statement. These include Applied Health Economics and Health Policy, Health and Quality of Life Outcomes, International Journal of Technology Assessment in Health Care, Journal of Medical Economics, Medical Decision Making, PharmacoEconomics and Quality of Life Research. We encourage other journals and 988 S. Petrou et al.

Table 1 Checklist of items to include when reporting a mapping study

Section/topic	Item number	Recommendation	Reported on page number/line number
Title and abstract			
Title	1	Identify the report as a study mapping between outcome measures. State the source measure(s) and generic, preference-based target measure(s) used in the study	
Abstract	2	Provide a structured abstract including, as applicable: objectives; methods, including data sources and their key characteristics, outcome measures used and estimation and validation strategies; results, including indicators of model performance; conclusions; and implications of key findings	
Introduction			
Study rationale	3	Describe the rationale for the mapping study in the context of the broader evidence base	
Study objective	4	Specify the research question with reference to the source and target measures used and the disease or population context of the study	
Methods			
Estimation sample	5	Describe how the estimation sample was identified, why it was selected, the methods of recruitment and data collection, and its location(s) or setting(s)	
External validation sample	6	If an external validation sample was used, the rationale for selection, the methods of recruitment and data collection, and its location(s) or setting(s) should be described	
Source and target measures	7	Describe the source and target measures and the methods by which they were applied in the mapping study	
Exploratory data analysis	8	Describe the methods used to assess the degree of conceptual overlap between the source and target measures	
Missing data	9	State how much data were missing and how missing data were handled in the sample(s) used for the analyses	
Modelling approaches	10	Describe and justify the statistical model(s) used to develop the mapping algorithm	
Estimation of predicted scores or utilities	11	Describe how predicted scores or utilities are estimated for each model specification	
Validation methods	12	Describe and justify the methods used to validate the mapping algorithm	
Measures of model performance	13	State and justify the measure(s) of model performance that determine the choice of the preferred model(s) and describe how these measures were estimated and applied	
Results			
Final sample size(s)	14	State the size of the estimation sample and any validation sample(s) used in the analyses (including both number of individuals and number of observations)	
Descriptive information	15	Describe the characteristics of individuals in the sample(s) (or refer back to previous publications giving such information). Provide summary scores for source and target measures, and summarise results of analyses used to assess overlap between the source and target measures	
Model selection	16	State which model(s) is(are) preferred and justify why this(these) model(s) was(were) chosen	
Model coefficients	17	Provide all model coefficients and standard errors for the selected model(s). Provide clear guidance on how a user can calculate utility scores based on the outputs of the selected model(s)	
Uncertainty	18	Report information that enables users to estimate standard errors around mean utility predictions and individual-level variability	
Model performance and face validity	19	Present results of model performance, such as measures of prediction accuracy and fit statistics for the selected model(s) in a table or in the text. Provide an assessment of face validity of the selected model(s)	
Discussion			
Comparisons with previous studies	20	Report details of previously published studies developing mapping algorithms between the same source and target measures and describe differences between the algorithms, in terms of model performance, predictions and coefficients, if applicable	
Study limitations	21	Outline the potential limitations of the mapping algorithm	

The MAPS Statement 989

Table 1 continued

Section/topic	Item number	Recommendation	Reported on page number/line number
Scope of applications	22	Outline the clinical and research settings in which the mapping algorithm could be used	
Other			
Additional information	23	Describe the source(s) of funding and non-monetary support for the study, and the role of the funder(s) in its design, conduct and report. Report any conflicts of interest surrounding the roles of authors and funders	

research interest groups to endorse the MAPS statement and authors to adhere to its principles.

4 The MAPS Explanation and Elaboration Paper

In addition to the MAPS reporting statement, we have produced a supporting Explanation and Elaboration paper [16] modelled on those developed for other reporting guidelines [9–14]. The reporting items contained within the MAPS statement are best understood by referring to the information contained within this accompanying document. The Explanation and Elaboration paper provides exemplars of good reporting practice identified from the published literature for each reporting item. In addition, it provides a detailed explanation to accompany each recommendation, supported by a rationale and relevant evidence where available. The development of the Explanation and Elaboration paper was completed following several iterations produced by members of the working group, after which the examples and explanations were shared with the Delphi panellists for final revisions to improve readability and their approval. The Explanation and Elaboration paper also summarises the characteristics of the Delphi panellists and provides detailed statistics for item ratings at each Delphi round.

5 Discussion

Over recent years, there has been a rapid increase in the publication of studies that use mapping techniques to predict health utility values. One recent review article identified 90 studies published up to the year 2013 reporting 121 mapping algorithms between clinical or health-related quality-of-life measures and the EQ-5D [2]. That review article excluded mapping algorithms targeted at other generic preference-based outcome measures that can generate health utilities, such as the SF-6D [17] and the Health Utilities Index [18], which have been the target of numerous other mapping algorithms (e.g. [1, 19–24]).

Moreover, the popularity of the mapping approach for estimating health utilities is unlikely to wane given the numerous contexts within health economic evaluation where primary data collection is challenging. However, mapping introduces additional uncertainty and collection of primary data with the preferred utility instrument is preferable.

The MAPS statement was developed to provide recommendations, in the form of a checklist of essential items, which authors should consider when reporting mapping studies. Guidance for the reporting of mapping studies was not previously available in the literature. The overall aim of the MAPS statement is to promote clarity, transparency and completeness of reporting of mapping studies. It is not intended to act as a methodological guide, nor as a tool for assessing the quality of study methodology. Rather, it aims to avoid misleading conclusions being drawn by readers, and ultimately policy makers, as a result of sub-optimal reporting. In keeping with other recent health research reporting guidelines, we have also produced an accompanying Explanation and Elaboration paper [16] to facilitate a deeper understanding of the 23 items contained within the MAPS statement. That paper should hopefully act as a pedagogical framework for researchers reporting mapping studies.

The development of the MAPS statement, and its Explanation and Elaboration document, was framed by recently published guidance for health research reporting guidelines [5]. The Delphi panel was composed of a multidisciplinary multi-national team of content experts and journal editors. The panel members included people experienced in conducting mapping studies; of the 84 researchers who were first authors on papers included in a recent review of EQ-5D mapping studies [2], 31 (36.9 %) were included as panellists. We have no evidence to believe that a larger panel would have altered the final set of recommendations. The Delphi methodologies that we applied included analytical approaches only recently adopted by developers of health reporting guidelines [15]. We are unable to assess whether a strict adherence to the MAPS checklist will increase the word counts of mapping reports. It is our view that the increasing use of online appendices by journals should permit comprehensive reporting even in the context of strict word limits for the main body of reports.

Evidence for other health research reporting guidelines suggests that reporting quality improved after the introduction of reporting checklists [25-27], although there is currently no empirical evidence that adoption of the MAPS statement will improve the quality of reporting of mapping research. Future research planned by the MAPS working group will include a before and after evaluation of the benefits (and indeed possible adverse effects) of the introduction of the MAPS statement. It will also be necessary to update the MAPS statement in the future to address conceptual, methodological and practical advances in the field. Potential methodological advances that might be reflected in an update might include shifts toward more complex model specifications, better methods for dealing with uncertainty and guidance on appropriate use of measures of prediction accuracy, such as mean absolute error and mean square error. The MAPS working group plans to assess the need for an update of the reporting checklist in 5 years' time.

6 Conclusion

This paper summarises a new reporting statement developed for studies that map onto generic preference-based outcome measures. We encourage health economics and quality-of-life journals to endorse the MAPS statement, promote its use in peer review, and update their editorial requirements and 'Instructions to Authors' accordingly.

Acknowledgments We are grateful to the following individuals who consented to being acknowledged publicly for their comments and participation in the Delphi Panel: Roisin Adams, Roberta Ara, Nick Bansback, Garry Barton, Andy Briggs, Chris Carswell, Gang Chen, Doug Coyle, Ralph Crott, Richard Edlin, Alastair Fischer, Juan Manuel Ramos-Goñi, Ola Ghatnekar, Patrick Gillard, Ning Yan Gu, Annie Hawton, Haomiao Jia, Kamran Khan, Billingsley Kaambwa, Nick Kontodimopoulos, Quang Le, Jason Madan, Katherine Mac-Gilchrist, David Meads, Duncan Mortimer, Bhash Naidoo, Angus Ng, Nalin Payakachat, Simon Pickard, Rafael Pinedo-Villanueva, Eleanor Pullenayegum, Jeff Richardson, Janet Robertson, Jeff Round, Donna Rowen, Sean Rundell, Paulos Teckle, Matthijs Versteegh, Feng Xie and Tracey Young.

This article is a joint publication by Applied Health Economics and Health Policy, Health and Quality of Life Outcomes, International Journal of Technology Assessment in Health Care, Journal of Medical Economics, Medical Decision Making, Pharmacoeconomics, and Quality of Life Research.

Compliance with Ethical Standards

No external funding was received for this study. All members of the working group acted on a voluntary basis.

RF is a shareholder and a director of a company (Clinvivo Limited) that provides services for Delphi studies. However, these services were not used in this study, and thus notwithstanding the connection, RF declares that he has no conflicts of interest. The remaining authors (SP, ORA, HD, LL, MO, AG) have no conflicts of interest to declare. All authors contributed to the study design and conduct, interpretation of results and report writing. SP acts as the guarantor.

Open Access This article is distributed under the terms of the Creative Commons Attribution-NonCommercial 4.0 International License (http://creativecommons.org/licenses/by-nc/4.0/), which permits any noncommercial use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made.

References

- Brazier JE, Yang Y, Tsuchiya A, Rowen DL. A review of studies mapping (or cross walking) non-preference based measures of health to generic preference-based measures. Eur J Health Econ. 2010;11(2):215–25. doi:10.1007/s10198-009-0168-z (1425919854).
- Dakin H. Review of studies mapping from quality of life or clinical measures to EQ-5D: an online database. Health Qual Life Outcomes. 2013;11:151. doi:10.1186/1477-7525-11-151 (1425925341).
- Chuang LH, Whitehead SJ. Mapping for economic evaluation. Br Med Bull. 2012;101:1–15. doi:10.1093/bmb/ldr049 (1427393724).
- Longworth L, Rowen D. NICE DSU Technical Support Document 10: the use of mapping methods to estimate health state utility values. Sheffield: Decision Support Unit, ScHARR, University of Sheffield; 2011.
- Moher D, Schulz KF, Simera I, Altman DG. Guidance for developers of health research reporting guidelines. PLoS Med. 2010;7(2):e1000217. doi:10.1371/journal.pmed.1000217 (1425988566).
- Moher D, Simera I, Schulz KF, Hoey J, Altman DG. Helping editors, peer reviewers and authors improve the clarity, completeness and transparency of reporting health research. BMC Med. 2008;6:13. doi:10.1186/1741-7015-6-13 (1425989043).
- Narahari SR, Ryan TJ, Aggithaya MG, Bose KS, Prasanna KS. Evidence-based approaches for the Ayurvedic traditional herbal formulations: toward an Ayurvedic CONSORT model. J Altern Complement Med. 2008;14(6):769–76. doi:10.1089/acm.2007. 0818 (1425997359).
- Mortimer D, Segal L, Sturm J. Can we derive an 'exchange rate' between descriptive and preference-based outcome measures for stroke? Results from the transfer to utility (TTU) technique. Health Qual Life Outcomes. 2009;7:33. doi:10.1186/1477-7525-7-33 (1426002631).
- Bossuyt PM, Reitsma JB, Bruns DE, Gatsonis CA, Glasziou PP, Irwig LM, et al. The STARD statement for reporting studies of diagnostic accuracy: explanation and elaboration. Ann Intern Med. 2003;138(1):W1–12 (1426070562).
- Tong A, Sainsbury P, Craig J. Consolidated criteria for reporting qualitative research (COREQ): a 32-item checklist for interviews and focus groups. Int J Qual Health Care. 2007;19(6):349–57. doi:10.1093/intqhc/mzm042 (1426070607).
- Vandenbroucke JP, von Elm E, Altman DG, Gotzsche PC, Mulrow CD, Pocock SJ, et al. Strengthening the Reporting of Observational Studies in Epidemiology (STROBE): explanation and elaboration. PLoS Med. 2007;4(10):e297. doi:10.1371/ journal.pmed.0040297 (1426070521).

The MAPS Statement 991

- Liberati A, Altman DG, Tetzlaff J, Mulrow C, Gotzsche PC, Ioannidis JP, et al. The PRISMA statement for reporting systematic reviews and meta-analyses of studies that evaluate health care interventions: explanation and elaboration. J Clin Epidemiol. 2009;62(10):e1-34. doi:10.1016/j.jclinepi.2009.06. 006 (1426070400).
- Moher D, Hopewell S, Schulz KF, Montori V, Gotzsche PC, Devereaux PJ, et al. CONSORT 2010 explanation and elaboration: updated guidelines for reporting parallel group randomised trials. BMJ. 2010;340:c869. doi:10.1136/bmj.c869 (1426070468).
- 14. Husereau D, Drummond M, Petrou S, Carswell C, Moher D, Greenberg D, et al. Consolidated Health Economic Evaluation Reporting Standards (CHEERS): explanation and elaboration. A report of the ISPOR Health Economic Evaluation Publication Guidelines Good Reporting Practices Task Force. Value Health. 2013;16(2):231–50. doi:10.1016/j.jval.2013.02.002 (1426070220).
- Fitch K, Bernstein S, Aguilar M, Burnand B, LaCalle J, Lazaro P, van het Loo M, McDonnell J, Vader J, Kahan J. The RAND/ UCLA appropriateness method user's manual. Santa Monica: RAND: 2001.
- Petrou S, Rivero-Arias O, Dakin H, Longworth L, Oppe M, Froud R, Gray A. The MAPS reporting statement for studies mapping onto generic preference-based outcome measures: explanation and elaboration. Pharmacoeconomics. 2015. doi:10.1007/s40273-015-0312-9.
- Brazier J, Roberts J, Deverill M. The estimation of a preferencebased measure of health from the SF-36. J Health Econ. 2002;21(2):271–92 (1426244550).
- Feeny D, Furlong W, Boyle M, Torrance GW. Multi-attribute health status classification systems. Health Utilities Index. Pharmacoeconomics. 1995;7(6):490–502 (1426244646).
- Yang Y, Wong MY, Lam CL, Wong CK. Improving the mapping of condition-specific health-related quality of life onto SF-6D

- score. Qual Life Res. 2014;23(8):2343–53. doi:10.1007/s11136-014-0668-x (1426244912).
- Lee L, Kaneva P, Latimer E, Feldman LS. Mapping the Gastrointestinal Quality of Life Index to short-form 6D utility scores.
 J Surg Res. 2014;186(1):135–41. doi:10.1016/j.jss.2013.08.006 (1426244970).
- Roset M, Badia X, Forsythe A, Webb SM. Mapping CushingQoL scores onto SF-6D utility values in patients with Cushing's syndrome. Patient. 2013;6(2):103–11. doi:10.1007/s40271-013-0010-7 (1426245033).
- Payakachat N, Tilford JM, Kuhlthau KA, van Exel NJ, Kovacs E, Bellando J, et al. Predicting health utilities for children with autism spectrum disorders. Autism Res. 2014;7(6):649–63. doi:10.1002/aur.1409 (1426245111).
- Goldfeld KS, Hamel MB, Mitchell SL. Mapping health status measures to a utility measure in a study of nursing home residents with advanced dementia. Med Care. 2012;50(5):446–51. doi:10. 1097/MLR.0b013e3182407e0d (1426245177).
- Chen G, McKie J, Khan MA, Richardson JR. Deriving health utilities from the MacNew Heart Disease Quality of Life Questionnaire. Eur J Cardiovasc Nurs. 2014;. doi:10.1177/ 1474515114536096 (1426245245).
- Moher D, Jones A, Lepage L, Group C. Use of the CONSORT statement and quality of reports of randomized trials: a comparative before-and-after evaluation. JAMA. 2001;285(15):1992–5 (1426256489).
- Delaney A, Bagshaw SM, Ferland A, Manns B, Laupland KB, Doig CJ. A systematic evaluation of the quality of meta-analyses in the critical care literature. Crit Care. 2005;9(5):R575–82. doi:10.1186/cc3803 (1426256559).
- Plint AC, Moher D, Morrison A, Schulz K, Altman DG, Hill C, et al. Does the CONSORT checklist improve the quality of reports of randomised controlled trials? A systematic review. Med J Aust. 2006;185(5):263–7 (1426256647).