

Appendix 3 (as supplied by the authors): Supplementary results

Table 1: Risk of confounding and characteristics of the outcome assessment

Trial	No risk of confounding ¹					Characteristics of the outcome assessment ²		
	Short time between assessments	Exactly same patients	Same type of assessors	Same type of assessment procedure	Effective blinding	Outcome subjectivity	Trial involvement of nonblind assessor	Vulnerability of outcome to nonblind patients
Cohen 2004 (1,2)	Yes	No	No	No	Yes	5	5	1
Oesterle 2000 (3)	Yes	Yes	Yes	No	No	5	4	5
Powell 2001 (4)	Yes	No	Yes	No	Yes	5	1	1
Burkhoff 1999 (5)	Yes	Yes	No	No	No	5	4	5
Wedekind 2006 (6)	Yes	No	Yes	Yes	No	5	5	5
Weaver 2009 (7)	Yes	Yes	Yes	Yes	Yes	5	5	4
Noseworthy 1994 (8)	Yes	Yes	Yes	Yes	No	5	5	5
Narins 2010 (9)	Yes	No	No	No	Yes	3	5	1
Ulm 1999 (10)	Yes	Yes	Yes	No	Yes	5	4	1
Meltzer 2003 (11,12)	Yes	No	Yes	Yes	No	5	5	5
Miller 2003 (13)	Yes	No	Yes	No	Yes	5	5	2
Taber 1983 (14)	Yes	No	Yes	Yes	Yes	3	5	3
Hylaform 2004 (15)	Yes	No	No	No	Yes	5	5	1
Landsman 2010 (16)	Yes	No	No	No	Yes	5	3	1
Iglesia 2010 (17)	Yes	Yes	No	Yes	Yes	3	5	1
Reddihough 2004 (18)	Yes	No	Yes	No	Yes	4	4	3

¹A "Yes" indicates that the trial is considered free from the potential confounding factor in question. Evaluations were conducted by authors masked to the comparison between blind vs. nonblind trial results (except for the evaluation of whether the patients in both type of assessments were exactly the same). ²Masked evaluation on 1-5 Likert scales (5 signify a high degree of, e.g. outcome subjectivity).

Table 2: Correlation and inter-observer agreement

Trials with blind & nonblind assessment of a measurement scale outcome			Studies validating the scale used in the trials ¹		
Trial	Outcome	Correlation blind/nonblind	Inter-observer agreement		Characteristics of the scale validation study
		rho ²	Weighted Kappa ³	ICC ⁴	
Cohen 2004 (1,2)	Facial Fold Assessment Scale (0-5)	0.50	na ⁵	0.89 ⁶	8 observers assessed 80 photographs of persons with facial wrinkles (19)
Oesterle 2000 (3)	CCSA ⁷ class (I-IV)	0.58	0.60	na	2 physicians assessed 75 patients referred for treadmill exercise tolerance testing (20)
Powell 2001 (4)	Nasal obstruction (10 cm VAS ⁸)	na	na		
Burkhoff 1999 (5)	CCSA ⁶ class (I-IV)	0.54	0.60	na	2 physicians assessed 75 patients referred for treadmill exercise tolerance testing (20)
Wedekind 2006 (6)	PAS ⁹ (0-52)	0.91	(0.78) ¹⁰		2 psychiatrists assessed 23 patients with panic disorder in out-patient unit (21)
Weaver 2009 (7)	UPDRS III ¹¹ (0-108)	na	na	0.82	2 neurologists assessed 24 patients with Parkinson's in community-based study (22)
Nosworthy 1994 (8)	EDSS ¹² (0-10)	0.89	na	0.78	2 neurologists assessed 125 patients with multiple sclerosis (23)
Narins 2010 (9)	Wrinkle Severity Rating Scale (0-4)	na	0.75	na	5 observers assessed 30 photographs of persons with facial wrinkles (24)
Ulm 1999 (10)	UPDRS III ¹¹ (0-108)	na	na	0.82	2 neurologists assessed 24 patients with Parkinson's in community-based study (22)
Meltzer 2003 (11,12)	CGI-SS ¹³	na	na		
Taber 1983 (14)	Illness severity score (0-3), day 1	na			
Miller 2003 (13)	Synechia (0-3), last follow-up	na			
Hylaform 2004 (15)	Severity Grading Scale (0-5)	na	0.85	na	13 observers assessed 32 persons with nasolabial folds (25)
Landsman 2010 (16)	Clinical assessment scale (0-3)	na	na		
Iglesia 2010 (17)	POP-Q ¹⁴ (0-IV)	0.67	0.64	na	2 observers assessed 45 women attending outpatient (uro)gynaecology clinics (26)
Reddihough 2004 (18)	GMFM ¹⁵ (0-264)	0.89 ¹⁶	na	0.99	2 observers assessed 26 children with cerebral palsy (27)

¹The identification of the validation studies was based on a literature search on PubMed, and was not the result of a systematic review. ²Spearman's correlation coefficient. ³Weighted Kappa or mean Weighted Kappa (confidence intervals not accessible); ⁴Intra-class correlation coefficient (95% confidence intervals only accessible in Reddihough 2004: (0.97-0.99)); ⁵not accessible; ⁶Blind outcome assessors in Cohen 2004 were reported to have ICC > 0.80; ⁷Canadian cardiovascular society (grading of) angina pectoris; ⁸Visual analogue scale; ⁹Panic and agoraphobia scale; ¹⁰Unclear what measure was used; ¹¹Unified Parkinson's Disease Rating Scale; ¹²Expanded disability status scale; ¹³Clinical global impression on suicide severity scale (7 point version); ¹⁴Pelvic organ prolapse quantification exam; ¹⁵Gross Motor Function Measure; ¹⁶Concordance correlation coefficient ranged from 0.78 to 0.99. We assumed that the nearly identical Spearman's correlation coefficient was 0.89.

Qualitative summary of results in trials with incomplete or unclear data:

Eight included trials had incomplete or unclear outcome data. Qualitative information, or results from other similar trials, indicated notable observer bias in three of these trials. Ash 1998 (28) was a split-body trial comparing the effect of tretinoin vs. L-ascorbic acid on 10 patients with striae alba. Assessment of percentage improvement was conducted by both blind and nonblind assessor. The means of the improvements were reported, and showed a clear difference favouring the nonblind assessment, but SD was not reported. We obtained a SD from a somewhat related trial (29), but it was debatable whether the trials were truly comparable. However, based on the SD, we derived at a tentative estimate dSMD of -0.81, indicating a substantial degree of observer bias.

Purdue 1997 (30) was a split-body trial comparing biosynthetic skin replacement vs. cadaver skin for burn wounds in 66 patients. Wound healing (percentage "take of autograft") was assessed by blind and nonblind assessors. The mean of the percentage take was reported inconsistently for the nonblind assessment, and SDs were not reported, so we decided to regard the data as too uncertain for inclusion in our main analysis. Based on the most conservative assessment and SD from a similar trial (31), we derived at a tentative dSMD of -0.48, indicating a substantial degree of observer bias.

Kadish 2011 (32) was a parallel group trial comparing cardiac contractility modulation vs. standard treatment in 428 patients with advanced heart failure. Assessment of NYHA class was conducted both by blind and nonblind assessors. The trial publication did not report the mean values of blind vs. nonblind assessments, and the company was unwilling to share the data. However, one of the authors, Dr. Daniel Burkhoff, informed us that there was a marked difference between effect estimates based on blind and nonblind assessment, similar to that of the two other cardiological trials included (3,5), with a pooled dSMD -0.56, indicating a substantial degree of observer bias.

There was indication of no or little bias in two trials. Bauman 2007 (33) was a large parallel group trial comparing three types of hyaluronic acid dermal fillers vs. collagen in 439 patients. Assessments of nasiolabel folds on the wrinkle assessment scale were conducted by blind and nonblind observers. Incomplete data based on two of the three intervention groups (277 patients) provided a tentative dSMD of -0.06, indicating low degree of observer bias.

Herberger 2011 (34) was a parallel group trial comparing the effect of ultrasound-assisted wound treatment vs. surgical debridement in 67 patients. Assessment of four dimensions of wound status was conducted by blind and nonblind assessors on a five point response scale. The authors provided us with individual patient data. We were unable to reproduce the published blinded results and number of patients was not identical. Further contact to the authors did not resolve the issue, so we decided to regard the data as too uncertain for inclusion in our main analysis. Our tentative estimate of dSMD was 0.12, indicating bias in the reversed direction.

In the remaining three trials there was no information on how disagreements between blind and nonblind assessors affected estimated intervention effects. Realmuto 1984 (35) was a parallel group trial comparing the effect of thiothixene vs. thioridazine in 21 schizophrenic adolescents. The blind and nonblind ratings on the Brief Psychiatric Rating Scale and Clinical Global Impression Scale "did not show statistically significant differences". Havel 1999 (36) was a parallel group trial comparing the effect of propofol vs. midazolam for procedural sedation in 91 patients. The weighted kappa value between the blind and nonblind ratings of Ramsey Sedation scores was reported as 0.74. Alam 2006 (37) was a split-body trial comparing the effect of single pulse dye laser treatment v no-treatment for scar appearance in 20 patients. The ratings of the blind and nonblind assessors of overall scar appearance were "highly correlated".

Conversion of dSMD to ratio of odds ratios (ROR)

To facilitate the comparison between the degree of observer bias in trials with measurement scale outcomes and binary outcomes we converted all SMDs to ORs, and subsequently all dSMDs to ROR. This conversion is based on assumptions of equal variance and logistic distributions of the measurements in each group (38).

The pooled result of all 16 trials produced an I^2 of 46% ($P = 0.02$), and a ratio of odds ratio of 0.66 (0.48 to 0.90).

An ROR < 1 indicates that nonblind outcome assessors generate a more optimistic estimate of the treatment effect than blind outcome assessors.

The result is coherent with that of our previous analyses of observer bias in 21 trials with binary outcomes, providing a pooled ratio of odds ratios of 0.64 (0.43 to 0.96) (39).

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