

1 **Professional medical writing support and the quality, ethics and timeliness**
2 **of clinical trial reporting: a systematic review**

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21 **Abstract**

22 **Background:** Many authors choose to work with professional medical writers when reporting the
23 results of clinical trials. We conducted a systematic review to examine the relationship between
24 professional medical writing support (PMWS) and the quality, ethics and timeliness of publications
25 reporting clinical trials.

26 **Methods:** Using terms related to ‘medical writer’ and ‘observational study’, we searched MEDLINE
27 and Embase (no date limits), as well as abstracts and posters from meetings of the International
28 Society for Medical Publication Professionals (ISMPP; 2014–2017). We also hand-searched the
29 journals *Medical Writing* and *The Write Stuff* (2014–2017), and the bibliographies of studies
30 identified in the electronic searches. We screened the results to identify studies that compared the
31 quality, ethics and timeliness of clinical trial publications written with and without declared PMWS.

32 **Results:** Our searches identified 97 potentially relevant studies, of which 89 were excluded during
33 screening and full paper review. The remaining eight studies compared 849 publications with PMWS
34 with 2073 articles developed without such support. In these eight studies, PMWS was shown to be
35 associated with: increased adherence to Consolidated Standards of Reporting Trials (CONSORT)
36 guidelines (in 3/3 studies in which this was assessed); publication in journals with an impact factor
37 (one study); a higher quality of written English (one study); and a lower likelihood of reporting non-
38 pre-specified outcomes (one study). PMWS was not associated with increased adherence to
39 CONSORT for Abstracts guidelines (one study) or with the impact of published articles (mean
40 number of citations per year, mean number of article views per year and Altmetric score; one study).
41 In studies that assessed timeliness of publication, PMWS was associated with a reduced time from last
42 patient visit in clinical trials to primary publication (one study), whereas time from submission to
43 acceptance showed inconsistent results (two studies).

44 **Conclusions:** This systematic review of eight observational studies suggests that PMWS increases the
45 overall quality of reporting of clinical trials and may improve the timeliness of publication.

46 **Keywords:** medical writing, medical writer, clinical trials transparency, reporting guidelines,
47 adherence

48 **Background**

49 Timely and complete reporting of the results of clinical trials is an ethical imperative [1]; it helps to
50 eliminate duplicative effort, enables researchers to develop more up-to-date study hypotheses and
51 allows clinicians and patients to judge the benefits or risks of different therapies. Although the
52 pharmaceutical industry has made great strides to address criticism for a perceived lack of
53 transparency in the disclosure of clinical trial results, the quality, ethics and timeliness of clinical trial
54 reporting remain closely scrutinized for both industry-funded and academically funded trials [2-6].

55 Pharmaceutical companies often offer authors professional medical writing support (PMWS) to assist
56 in the reporting of clinical trial results [7]. International guidelines endorse the acknowledgement of
57 PMWS [8,9], and the proportion of articles in the medical literature with such an acknowledgement is
58 6–19% [7,10,11]. We conducted a systematic review to identify and analyse published studies that
59 investigated the association between PMWS and the quality, ethics and timeliness of clinical trial
60 reporting.

61 **Methods**

62 **Systematic literature search**

63 Published studies relating to medical writing were identified through a systematic literature review.
64 Cochrane, Embase, MEDLINE In-Process & Other Non-Indexed Citations, and MEDLINE 1946–
65 present were searched on 8 March 2018 via the Ovid platform.

66 The search strategy comprised terms relating to medical writing, medical publication professional and
67 medical communication, and was combined with terms for observational, cross-sectional or

68 epidemiological studies, with no limits on date, language or country in which the research was
69 conducted (Figure 1).

70 **Supplementary searches**

71 Supplementary searches were conducted of the International Society for Medical Publication
72 Professionals (ISMPP) congress proceedings (which are published as supplementary articles in
73 *Current Medical Research Opinion*), and the journals *Medical Writing* and *The Write Stuff* (which are
74 available via the European Medical Writers Association [EMWA] website) using the terms
75 ‘medical writ*’ and ‘medical publication professional’. Supplementary searches were limited to
76 2014–2017. We contacted the corresponding authors of congress abstracts identified in the
77 supplementary searches to request access to full posters/presentations. The bibliographies of studies
78 identified in the electronic searches were also reviewed to identify additional relevant references.

79 **Study selection and data collection**

80 All identified studies were screened against inclusion and exclusion criteria in accordance with the
81 2009 Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines
82 [12]. For congress abstracts identified in the supplementary searches, full posters were obtained from
83 the ISMPP website or from the authors. Identified congress abstracts were excluded as ‘duplicates’ if
84 a full version of the study had been published. Studies eligible for inclusion were in English, and
85 compared the quality, ethics or timeliness of articles reporting clinical trials that had been developed
86 with and without acknowledged PMWS. Studies that did not directly compare clinical trial
87 publications that had been developed with and without PMWS were excluded, as were those that
88 reported outcomes that were unrelated to quality, ethics or timeliness, and those that assessed study
89 types other than clinical trials.

90 Details of study methodology, study size, main outcome measures, quality-related outcomes
91 (e.g. adherence to Consolidated Standards of Reporting Trials [CONSORT] or CONSORT for
92 Abstracts [CONSORT-A]), ethics-related outcomes (e.g. reporting of non-pre-specified outcomes)
93 and timeliness-related outcomes were extracted from each eligible study. The influence of PMWS

94 was classified as positive, non-significant or negative for each study, based on the results and
95 statistical analyses reported in each publication.

96 **Results**

97 **Search results**

98 Our searches identified 75 potentially relevant publications after exclusion of 22 duplicate
99 publications; 70 were excluded during screening and full paper review, and three were identified in
100 bibliographies of identified studies (Figure 1). Of the eight included studies, three were full
101 publications (two in peer-reviewed journals [13,14], one in a non-peer reviewed journal [15]), and
102 five were congress abstracts (four poster presentations [16-19], one oral presentation [20]). Although
103 no date limit was included in the search strategy, only two of the identified studies were published
104 before 2015: one in 2006 [7] and the other in 2010 [15] (Table 1). The eight included studies analysed
105 849 articles that had been developed with PMWS and 2073 articles developed without.

106 **Quality of reporting**

107 Of the identified studies comparing articles developed with and without PMWS, three assessed
108 adherence to CONSORT guidelines [14,15,19]. Each of these studies, using a different statistical
109 approach to assess adherence, showed that PMWS was associated with increased adherence to
110 CONSORT guidelines (Table 2). Articles developed with PMWS were significantly more likely to
111 report completely at least 50% of the assessed CONSORT items ($p < 0.05$) [14,21] and to comply
112 with more CONSORT items than articles without PMWS ($p < 0.05$) [15]. Similarly, articles with 80–
113 100% compliance with CONSORT items were significantly more likely to have been developed with
114 PMWS than those with less than 80% compliance ($p < 0.0001$) [19]. Looking at individual
115 CONSORT items, one identified study showed that articles with PMWS were significantly more
116 likely to report all important adverse events or side effects than those without PMWS [15], and
117 another showed that PMWS increased adherence to six of 12 CONSORT items assessed: specification
118 of primary outcome; sample size calculation; type of randomization; publication of a participant flow
119 diagram; provision of dates defining recruitment and follow-up; and details of trial registration [14].
120 Additionally, in this study, another CONSORT item (who generated the allocation sequence) was

121 only reported in 5/110 articles developed with PMWS and none of the 123 articles without PMWS;
122 thus, a relative risk could not be calculated [14]. One additional study assessed adherence to
123 CONSORT-A and showed that PMWS was not associated with an overall increase in adherence [13];
124 PMWS was associated with lower levels of adherence with respect to reporting of study setting and
125 higher levels of adherence in relation to disclosure of harms/side effects and funding source in the
126 abstract [13].

127 Two studies which represented different analyses of the same group of articles looked at other
128 markers of quality in reporting (Table 2) [14,17]. In these studies, PMWS was positively associated
129 with various measures of reporting quality, including a higher standard of written English ($p < 0.01$)
130 [14,21], higher likelihood of publication in a journal with an impact factor ($p = 0.001$) [17], and
131 higher mean impact factor of the journal accepting the article ($p < 0.001$) [17]. However, there was no
132 association between PMWS and article-level measures of impact, such as mean number of citations
133 per year ($p = 0.11$), mean number of article views per year ($p = 0.84$) and Altmetric score ($p = 0.55$)
134 (Table 2) [17].

135 **Ethics of publication**

136 Of the identified studies, one examined the relationship between outcome reporting and PMWS using
137 data from the publicly available Centre for Evidence-Based Medicine Outcome Monitoring Project
138 (COMParE) [22]. PMWS was associated with the reporting of fewer non-pre-specified outcomes
139 ($p = 0.028$) [16].

140 **Timeliness of publication**

141 Three studies looked at the timeliness of clinical trial reporting in articles developed with or without
142 PMWS (Table 2) [14,18,20]. The only study investigating the complete manuscript development time,
143 from last patient visit in clinical trials to article publication, showed that PMWS was associated with
144 reduced time to publication [18]. Two studies investigating the timing of one step in the process, from
145 manuscript submission to acceptance, showed inconsistent results [14,20]. In the first of these studies,
146 PMWS was associated with increased time from manuscript submission to acceptance, although the

147 mean number of versions submitted was unchanged [14]; in the second study, time from manuscript
148 submission to acceptance was reduced, but not significantly [20].

149 **Conclusions**

150 This systematic review aimed to identify and evaluate studies assessing the effects of PMWS on
151 quality, ethics and timeliness of clinical trial reporting. Overall findings from eight studies assessing
152 849 articles developed with PMWS and 2073 articles developed without PMWS suggest a positive
153 association between PMWS and improvements in clinical trial reporting. These results were
154 consistent across measures of quality (adherence to CONSORT guidelines and quality of written
155 English), ethics (reporting of non-pre-specified outcomes) and timeliness (time to publication). The
156 improvement in CONSORT adherence associated with PMWS is perhaps unsurprising, given that
157 professional medical writers are routinely trained in Good Publication Practice (GPP3) for the
158 development of peer-reviewed manuscripts [23]; GPP3 guidelines state that authors should follow
159 established reporting standards, including CONSORT [8,9]. Although PMWS was associated with
160 improved adherence to CONSORT, it was not associated with improved adherence to CONSORT-A,
161 suggesting that although professional medical writers improve disclosure overall, they may need to
162 prioritize improving the reporting in the abstract (which is all that is read by many readers).

163 The improvements in manuscript quality may not be reflected by increased article impact and social
164 media attention. In the one study identified in our systematic review, which examined measures of
165 article impact, there were no significant differences between articles developed with and without
166 PMWS in relation to Altmetric score, number of citations per year and number of article views per
167 year. Medical publications professions have no influence on the subject matter or relevance of the
168 clinical trial and, as such, PMWS may not be expected to affect an article's post-publication impact.

169 It is important that authors remain transparent about which specific clinical trial outcomes will be
170 measured and reported. The COMPare project determined the proportion of pre-specified and non-
171 pre-specified outcomes that were reported in clinical studies published in the top five medical journals
172 over a 3-month period [22]. In the present systematic review, one included study conducted a sub-

173 analysis of the publicly available COMPare data and assessed the relationship between PMWS and
174 outcome reporting. The authors found that fewer non-pre-specified outcomes were reported for
175 articles developed with PMWS than for those developed without. This is not the only study to have
176 shown a positive association between PMWS and publication ethics. For instance, a recent study
177 showed that PMWS is associated with increased transparency relating to the source of funding, the
178 author disclosures of financial interest and the acknowledgements of conflicts of interest (or lack
179 thereof) in health economics and outcomes research publications [24]; another study showed that, of
180 214 publications retracted owing to misconduct between January 1966 and February 2008, only three
181 declared PMWS [25].

182 One included study looking at the influence of PMWS on timeliness found that PMWS was associated
183 with reduced time from last patient visit to article publication. This period includes processes in which
184 professional medical writers are involved and have a major role, namely manuscript preparation,
185 editing and submission. Two other included studies that examined the influence of PMWS on time
186 from manuscript submission to acceptance revealed mixed results. One of the studies found that time
187 to acceptance was reduced with PMWS, but that the difference was not statistically significant. The
188 other study found that time to acceptance was increased with PMWS; however, it should be noted that
189 the period from submission to article acceptance is not primarily the responsibility of professional
190 medical writers.

191 Clinicians have reported lack of time as a common reason for non-publication of research findings
192 [26-28]. By specializing in preparation of clinical trial publications, professional medical writers are
193 well placed to aid in the rapid dissemination of trial findings under the direction of the authors,
194 subject to strict publication guidelines [9]. In fact, results from a recent survey showed that authors
195 who use PMWS were more likely to have published as first author at least once in the previous 2
196 years [29], suggesting that PMWS can also improve overall publication rates.

197 This systematic review has some limitations, notably that study inclusion was largely based on the
198 assumption that differences in outcomes were attributable to PMWS. It is possible that other factors

199 caused these differences in quality and timeliness. This issue may affect the results of individual
200 studies, but this systematic review combined results from different studies looking at different
201 outcomes of interest, and showed a consistent benefit of PMWS on manuscript quality (including
202 adherence to publication guidelines, quality of written English and publication in high-quality
203 journals), ethics (reporting of pre-specified outcomes) and timeliness (time from completion of trial to
204 publication). Taken together, the findings of this systematic review support the conclusion that
205 PMWS has a positive impact on the high-quality, ethical and timely dissemination of clinical trial
206 data.

207 The included studies classified articles as having been developed with PMWS only when there was a
208 clear acknowledgement of this support. As such, it is possible that some of the studies classified as
209 having been developed with no PMWS might have had PMWS but had simply failed to acknowledge
210 it. By classifying publications with no clear acknowledgment of PMWS as ‘without PMWS’, the
211 studies identified in this systematic review may have underestimated the effects of PMWS. To
212 minimize the risk of publication bias we employed a broad search strategy with no limits on date,
213 country, language or type of observational study. Most of the identified studies were sourced from
214 conference proceedings (for which the full poster or oral presentation was available in 4/5 cases) and
215 one was published in a non-peer-reviewed journal.

216 In the identified studies, the outcome measures chosen were widely accepted as measures of quality
217 and completeness. For instance, CONSORT is an independently developed measure of reporting
218 standards recommended by the International Committee of Medical Journal Editors and also medical
219 publications and medical writing societies, including ISMPP, EMWA and the American Medical
220 Writers Association [9]. Other outcomes of interest assessed in this review were assigned
221 independently of the investigators involved in each of the articles analysed in each included study
222 (e.g. standard of written English – assessed during peer review of analysed articles [17]). As such, in
223 this systematic review, we have been successful in analysing a range of outcomes assessed in
224 observational (‘real-world’) studies in a standardized manner that minimizes publication bias.

225 Further research is needed to elucidate the role of PMWS in clinical trial publication, particularly with
226 regard to productivity and added value [30]. Further research is also required to assess the impact of
227 PMWS in other types of studies published by the pharmaceutical industry, such as observational
228 studies and systematic reviews. As our systematic review identified that most studies of PMWS have
229 only been presented at conferences or published in non-peer-reviewed journals, it is crucial that future
230 studies are published in full in peer-reviewed journals.[31]

231 Currently, the pharmaceutical industry is more likely than non-industry institutions to disclose clinical
232 trial results properly [32]. This is probably due to a larger investment in internal processes and
233 infrastructure, which includes the use of professional medical writing support. In fact, there have been
234 calls for professional medical writers and publication experts to be employed by academic institutions
235 [33,34]. Additionally, in a survey looking at attitudes to PMWS, academic and clinician respondents
236 to an online survey were generally accepting of PMWS, particularly its influence on editing, journal
237 styling and adherence to reporting guidelines, with 84% of respondents stating that they valued
238 PMWS [35]. In this survey, 82.9% of respondents felt that it was acceptable to receive PMWS; in
239 another survey, PMWS was seen as ‘adding value to publication development’ by almost 90% of
240 participants [35]. Our systematic review appraising current research in this area helps to substantiate
241 the positive attitude to PMWS that is held by clinical and academic professionals seeking to ensure
242 the ethical, accurate and timely publication of clinical trials.

243 **List of abbreviations:** CI, confidence interval; COMPare, Centre for Evidence-Based Medicine
244 Outcome Monitoring Project; CONSORT, Consolidated Standards of Reporting Trials;
245 CONSORT-A, CONSORT for Abstracts; EMWA, European Medical Writers Association;
246 FDA, Food and Drug Administration; GPP, Good Publication Practice; IQR, interquartile range;
247 ISMPP, International Society for Medical Publication Professionals; OR, odds ratio;
248 PMWS, professional medical writing support; PRISMA, Preferred Reporting Items for Systematic
249 Reviews and Meta-Analyses; RCT, randomized controlled trial; RR, relative risk; SD, standard
250 deviation.

251 **Declarations:**

252 **Ethics approval and consent to participate**

253 Not applicable.

254 **Consent for publication**

255 Not applicable

256 **Availability of data and material**

257 Data sharing is not applicable to this article as no datasets were generated or analysed during the
258 current study.

259 **Competing interests**

260 Obaro Evuarherhe, Richard White and Christopher C Winchester are employees of Oxford
261 PharmaGenesis, Oxford, UK. William Gattrell is an employee of Ipsen Pharma, Milton Park, UK.
262 Christopher C Winchester and Richard White are directors of, and own shares in, Oxford
263 PharmaGenesis Holdings Ltd.

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266 **Authors' contributions**

267 The study was conceived by CW and drafted by OE, WG, RW and CW.

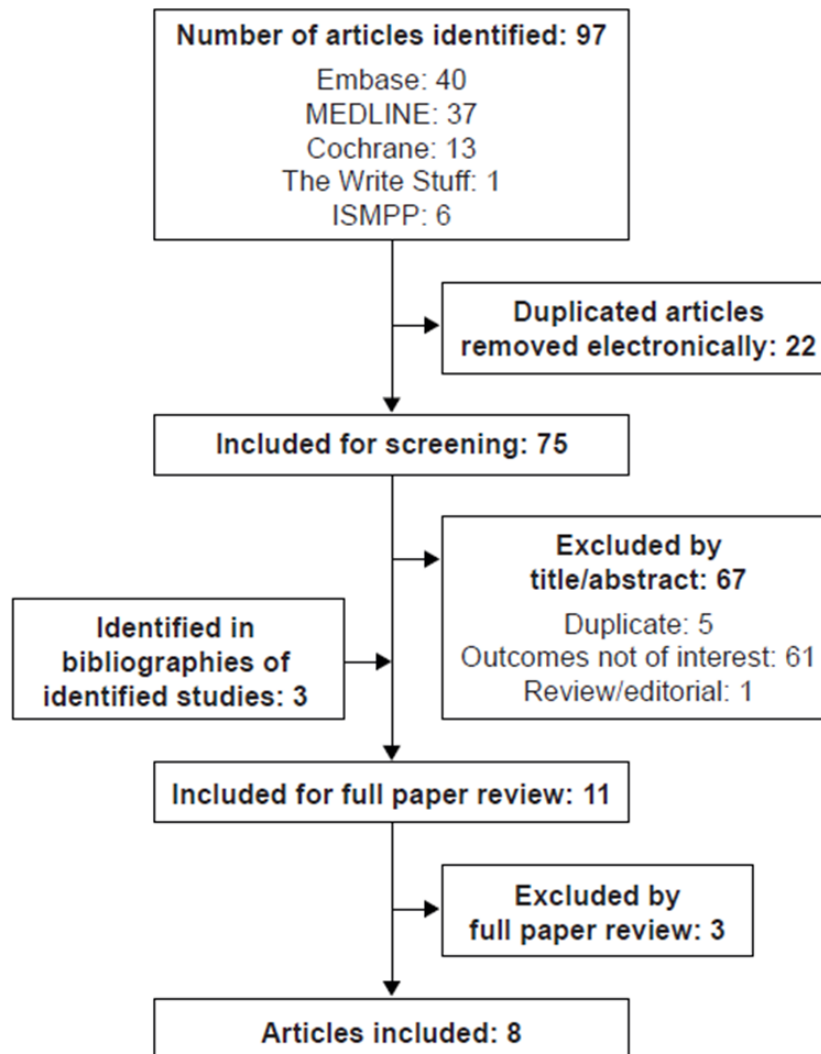
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269 Results of this systematic review were presented as a poster at the 2018 European Meeting of ISMPP
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272 poster for one of the identified congress abstracts.

273 **Tables and figure legend**

274 **Figure 1.** PRISMA diagram of included and excluded studies



Searches

- 1 (medical writer* or medical writing or medical publication professional* or medical communication or medcomms).mp.
- 2 ((observational adj (study or studies)) or (cross sectional adj (study or studies)) or (epidemiologic\$ adj (study or studies))).mp. or exp study/ or exp trial/
- 3 and/1-2

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276 ISMPP, International Society for Medical Publication Professionals; PRISMA, Preferred Reporting

277 Items for Systematic Reviews and Meta-Analyses.

Table 1. Overview of included studies

First author, year	Number of included studies		Publication type	Description of analysed articles
	With PMWS	Without PMWS		
Gattrell, 2016 [14]	110	123	Peer-reviewed publication	<ul style="list-style-type: none"> Articles reporting RCT results published in BioMed Central journals Biomed Central journals have been used in previous studies of adherence to CONSORT guidelines [37]
Gattrell, 2016 [17]	110	123	Poster presentation	<ul style="list-style-type: none"> Articles reporting RCT results published in BioMed Central journals (same cohort of articles analysed in Gattrell <i>et al.</i> [14])
Gattrell, 2017 [16]	17	49	Poster presentation	<ul style="list-style-type: none"> Sub-analysis of outcomes reported in the top five medical journals comparing each article with its corresponding study protocol or clinical trial registry entry using publicly available COMPare data The COMPare project is evaluating outcome reporting in clinical trials by comparing publications with the respective registry entries [22]
Jacobs, 2010 [15]	152	69	Non-peer-reviewed publication	<ul style="list-style-type: none"> RCTs published between October 2004 and August 2009 in the journal <i>Current Medical Research and Opinion</i>

				<ul style="list-style-type: none"> • <i>Current Medical Research and Opinion</i> almost exclusively publishes industry-funded studies
Mills, 2017 [13]	66	397	Peer-reviewed publication	<ul style="list-style-type: none"> • RCTs published between 2011 and 2014 in five high-impact medical journals: <i>The New England Journal of Medicine</i>, <i>Annals of Internal Medicine</i>, <i>The Lancet</i>, <i>The BMJ</i> and <i>JAMA</i> • All included articles had been analysed in a cross-sectional study examining reporting quality of RCTs [38]
Shah, 2015 [19]	40	103	Poster presentation	<ul style="list-style-type: none"> • Neuroscience and cardiology RCTs published between 2009 and 2014 in different journals from the Asia-Pacific region
Shah, 2016 [18]	404	392	Poster presentation	<ul style="list-style-type: none"> • RCTs conducted to gain US FDA approval in 2014 • Innovative and novel drugs and new molecules approved by the FDA in 2014, identified in FDA reports
Woolley, 2006 [7]	60	940	Congress abstract	<ul style="list-style-type: none"> • Original research articles published up to January 2005 from each of 10 high-impact factor, international, peer-reviewed medical journals from a range of therapeutic areas

COMPare, Centre for Evidence-Based Medicine Outcome Monitoring Project; CONSORT, Consolidated Standards of Reporting Trials; FDA, Food and Drug

Administration; PMWS, professional medical writing support; RCT, randomized controlled trial.

Table 2. Summary of results

First author, year	Outcome measured	Effect of PMWS		
		Positive	Non-significant	Negative
Gattrell, 2016 [14]	Adherence to CONSORT guidelines	The proportion of articles that completely reported at least 50% of the assessed CONSORT items <ul style="list-style-type: none"> • With PMWS: 43/110 articles (39.1%; 95% CI: 29.9–48.9) • Without PMWS: 26/123 articles (21.1%; 95% CI: 14.3–29.4) 		
Jacobs, 2010 [15]		Logistic regression analysis showed that CONSORT items were significantly more likely to be completed in papers with a clear acknowledgement of PMWS than in those without (OR 1.44; 95% CI: 1.04–2.00; $p = 0.03$)		
Shah, 2015 [19]		23/97 articles with PMWS (24%) had		

		80–100% CONSORT adherence, whereas 5/105 articles developed without PMWS (5%) had 80–100% CONSORT adherence ($p < 0.0001$)		
Mills, 2017 [13]	Adherence to CONSORT-A guidelines		The mean proportion of CONSORT-A items reported was similar with and without PMWS (64.3% vs 66.5%, respectively; $p = 0.30$); PMWS was associated with a lower level of compliance with reporting of study setting (RR 0.40; 95% CI: 0.23–0.70) and a higher level of adherence to disclosure of harms/side effects (RR 2.04; 95% CI: 1.37–3.03) and funding source (RR 1.75; 95% CI: 1.18–2.60)	

Gattrell, 2016 [14]	Quality of written English	<p>Proportion of articles rated by all reviewers during peer review as having an acceptable standard of written English</p> <ul style="list-style-type: none"> • With PMWS: 81.1% (43/53 articles; 95% CI: 67.6–90.1) • Without PMWS: 47.9% (23/48 articles; 95% CI: 33.5–62.7) 		
Gattrell, 2016 [17]	Publication in journal with an impact factor	Likelihood of publication in journal with an impact factor was significantly improved with PMWS ($p = 0.001$)		
	Mean impact factor of publication	Mean impact factor of publication was significantly improved with PMWS ($p < 0.001$)		

Gattrell, 2017 [16]	Reporting of non-pre-specified outcomes	Articles developed with PMWS reported fewer non-pre-specified outcomes than both industry-funded ($p = 0.028$) and non-industry-funded articles ($p < 0.01$) developed without PMWS		
Gattrell, 2016 [17]	Mean number of citations per year		Mean number of citations per year was not significantly improved with PMWS ($p = 0.11$)	
	Mean number of article views per year		Mean number of article views per year was not significantly improved with PMWS ($p = 0.84$)	
	Altmetric score		Altmetric score was not significantly improved with PMWS ($p = 0.55$)	

Gattrell, 2016 [14,21]	Manuscript acceptance time			Time from manuscript submission to acceptance was increased with PMWS (167 days [IQR 114.5–231 days] vs 136 days [IQR 77–193 days], $p < 0.01$); mean number of versions submitted was unchanged
Shah, 2016 [18]	Time to publication	Time to publication from last patient visit in clinical trials was reduced with PMWS (18.6 [SD 13.2] months vs 30.8 [SD 11.7] months)		
Woolley, 2006 [7]	Manuscript acceptance time		Time from manuscript submission to acceptance was reduced with PMWS (83.6 days vs 132.2 days), although this difference was not statistically significant ($p = 0.053$)	

CI, confidence interval; CONSORT, Consolidated Standards of Reporting Trials; CONSORT-A, CONSORT for Abstracts; IQR, interquartile range; OR, odds ratio;

PMWS, professional medical writing support; RR, relative risk; SD, standard deviation.

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