



OPEN ACCESS

# Effect size reporting among prominent health journals: a case study of odds ratios

Brian Chu <sup>1</sup>, Michael Liu,<sup>2</sup> Eric C Leas,<sup>3</sup> Benjamin M Althouse,<sup>4</sup> John W Ayers<sup>5</sup>

10.1136/bmjebm-2020-111569

<sup>1</sup>Perelman School of Medicine, University of Pennsylvania, Philadelphia, Pennsylvania, USA

<sup>2</sup>University of Oxford, Oxford, UK

<sup>3</sup>Department of Family Medicine and Public Health, Division of Health Policy, University of California San Diego, La Jolla, California, USA

<sup>4</sup>Epidemiology, Institute for Disease Modeling, Bellevue, Washington, USA

<sup>5</sup>Department of Medicine, Division of Infectious Diseases and Global Health, University of California San Diego, La Jolla, California, USA

Correspondence to: **Brian Chu**, University of Pennsylvania Perelman School of Medicine, Philadelphia, PA 19104, USA; [brianchu2010@gmail.com](mailto:brianchu2010@gmail.com)



© Author(s) (or their employer(s)) 2021. Re-use permitted under CC BY-NC. No commercial re-use. See rights and permissions. Published by BMJ.

**To cite:** Chu B, Liu M, Leas EC, *et al.* *BMJ Evidence-Based Medicine* 2021;**26**:184.

## ABSTRACT

**Background** The accuracy of statistical reporting that informs medical and public health practice has generated extensive debate, but no studies have evaluated the frequency or accuracy of effect size (the magnitude of change in outcome as a function of change in predictor) reporting in prominent health journals.

**Objective** To evaluate effect size reporting practices in prominent health journals using the case study of ORs.

**Design** Articles published in the American Journal of Public Health (AJPH), Journal of the American Medical Association (JAMA), New England Journal of Medicine (NEJM) and PLOS One from 1 January 2010 through 31 December 2019 mentioning the term ‘odds ratio’ in all searchable fields were obtained using PubMed. One hundred randomly selected articles that reported original research using ORs were sampled per journal for in-depth analysis.

**Main outcomes and measures** We report prevalence of articles using ORs, reporting effect sizes from ORs (reporting the magnitude of change in outcome as a function of change in predictor) and reporting correct effect sizes.

**Results** The proportion of articles using ORs in the past decade declined in JAMA and AJPH, remained similar in NEJM and increased in PLOS One, with 6124 articles in total. Twenty-four per cent (95% CI 20% to 28%) of articles reported the at least one effect size arising from an OR. Among articles reporting any effect size, 57% (95% CI 47% to 67%) did so incorrectly. Taken together, 10% (95% CI 7% to 13%) of articles included a correct effect size interpretation of an OR. Articles that used ORs in AJPH more frequently reported the effect size (36%, 95% CI 27% to 45%), when compared with NEJM (26%, 95% CI 17.5% to 34.7%), PLOS One (22%, 95% CI 13.9% to 30.2%) and JAMA (10%, 95% CI 3.9% to 16.0%), but the probability of a correct interpretation did not statistically differ between the four journals ( $\chi^2=0.56$ ,  $p=0.90$ ).

**Conclusions** Articles that used ORs in prominent journals frequently omitted presenting the effect size of their predictor variables. When reported, the presented effect size was usually incorrect. When used, ORs should be paired with accurate effect size interpretations. New editorial and research reporting standards to improve effect size reporting and its accuracy should be considered.

## Summary box

### What is already known about this subject?

▶ Although odds ratios (ORs) are frequently used, it is unknown whether ORs are reported correctly in peer-reviewed articles in prominent health journals.

### What are the new findings?

▶ In a random sample of 400 articles from four journals, reporting of ORs frequently omitted reporting effect sizes (magnitude of change in outcome as a function of change in predictor). When interpretations occurred, they were usually incorrect. Taken together, 10% (95% CI 7% to 13%) of articles correctly reported the effect size of an OR.

### How might it impact on clinical practice in the foreseeable future?

▶ Statistical results must be paired with accurate interpretations of effect size. Omitting effect size estimates poses challenges for decision-makers who synthesise inferences across articles. New editorial and research standards to encourage effect size reporting and accuracy in research using ORs should be considered.

## Introduction

In response to concerns about the accuracy of statistical reporting that informs medical and public health practice, standardised reporting guidelines (eg, Strengthening the Reporting of Observational Studies in Epidemiology (STROBE), Consolidated Standards of Reporting Trials, Preferred Reporting Items for Systematic Reviews and Meta-Analyses and others) have been developed.<sup>1</sup> Increasingly, describing the effect size (the magnitude of change in outcome as a function of change in predictor) is prioritised over reporting p-values.<sup>2,3</sup> Effect size forms the basis of both clinical importance (eg, treatments achieving statistically, but not clinically, significant improvements in outcomes are unlikely to be implemented) and practice guidelines (eg, treatments with larger effect sizes, with all else equal, are generally preferred). However, few studies have systematically quantified the

**Table 1** Criteria for labelling presence and correctness of interpretation of an OR

A *substantive effect size interpretation* (interpretation), for the purpose of this study, is a statement about the magnitude of change in the outcome as a function of change in the predictor from the reported OR. Solely reporting the OR or its directionality was not considered a substantive interpretation. A *correct substantive interpretation* is an interpretation that accurately reflects the definition of the OR, as described in commentary by Davies *et al*<sup>6</sup> and Norton *et al*.<sup>7</sup> For example, if the OR were 1.5, the interpretation “50% increase in odds” would be correct, while “50% more likely” would be incorrect. Correct interpretations could also include other interpretations resulting from logistic regression that use the OR as an intermediary (eg, change in probability, marginal risk).

Example phrases	Label	Reason
“... male sex was associated with seeking treatment (OR=2)...”	No interpretation	The OR was presented as a parenthetical statement only.
“... was associated with decreased odds (OR=0.5) ...”	No interpretation	Only the direction of the association was reported.
“... were three times more likely (OR=3) ...”	Incorrect interpretation	An interpretation was made by incorrectly expressing ‘odds’ as ‘likeliness’.
“... was associated with a 30% reduction in the log odds (OR=0.7) ...”	Incorrect interpretation	An interpretation was made by expressing the ratio of log odds but reported the OR.
“... were associated with a threefold increase in the odds (OR=3) ...”	Correct interpretation	An interpretation was made by expressing the ratio of odds.
“... each was associated with a 10% reduction in the odds of treatment failure (OR=0.90) ...”	Correct interpretation	An interpretation was made by expressing the ratio of odds.

accuracy of statistical reporting in practice,<sup>4 5</sup> and none have evaluated how articles in prominent health journals report the effect sizes of their results.

To this end, we conducted a case study of how effect sizes were reported among studies using ORs from prominent health journals. “Odds” are defined as the ratio of the probability of an outcome occurring to the probability of that outcome not occurring; an OR is the ratio of odds of an outcome between two groups. ORs do not easily translate into colloquial effect size interpretations, but numerous commentaries on how to interpret the magnitude of change in outcome as a function of change in predictor have been published.<sup>6-8</sup> As a result, we evaluated if articles published in prominent health journals using ORs reported the effect size of their results and whether these reports were accurate.

## Methods

### Eligible articles

Articles published in the *American Journal of Public Health* (AJPH), *Journal of the American Medical Association* (JAMA), *New England Journal of Medicine* (NEJM) and *PLOS One* from 1 January 2010 through 31 December 2019 mentioning “odds ratio” were obtained using PubMed by searching all available fields, including the title, abstract and keywords. These journals represent prominent medical (JAMA being the most circulated, NEJM having the highest impact factor) and public health (AJPH being published by the largest public health society) journals, and the highest-volume publisher (*PLOS One* publishing >200 000 articles in the past decade), and help set standards for the quality of scientific reporting in their domains. The search term “odds ratio” was

chosen because we sought to characterise statistical reporting in relation to ORs and alternative keywords provided low precision for discovering ORs (eg, only 31% of articles mentioning “odds ratio” also mentioned “logistic regression”). Next, three authors (BC, ML, JWA) randomly sampled articles to identify original research reports that included covariates. We excluded bivariable studies without covariates (nearly always trials) where authors may rely on descriptive results to portray effect size and use ORs as statistical tests rather than to estimate effect sizes. A quota of 100 articles was obtained per journal. Because only 94 NEJM articles met inclusion criteria, 6 articles were obtained from 2009.

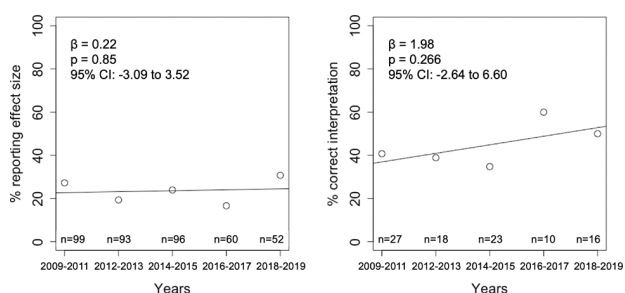
### Qualitative assessment of random samples

The same authors independently coded whether articles reported the effect sizes of ORs by describing the magnitude of change in the outcome as a function of change in the predictor ( $\kappa=0.90$ , overlapping sample  $n=28$ ). Solely reporting the OR or its directionality was insufficient (table 1). Both the results reported in the abstract and complete text were read to identify interpretations. Articles with at least one effect size interpretation were labelled accordingly, even if other ORs were uninterpreted.

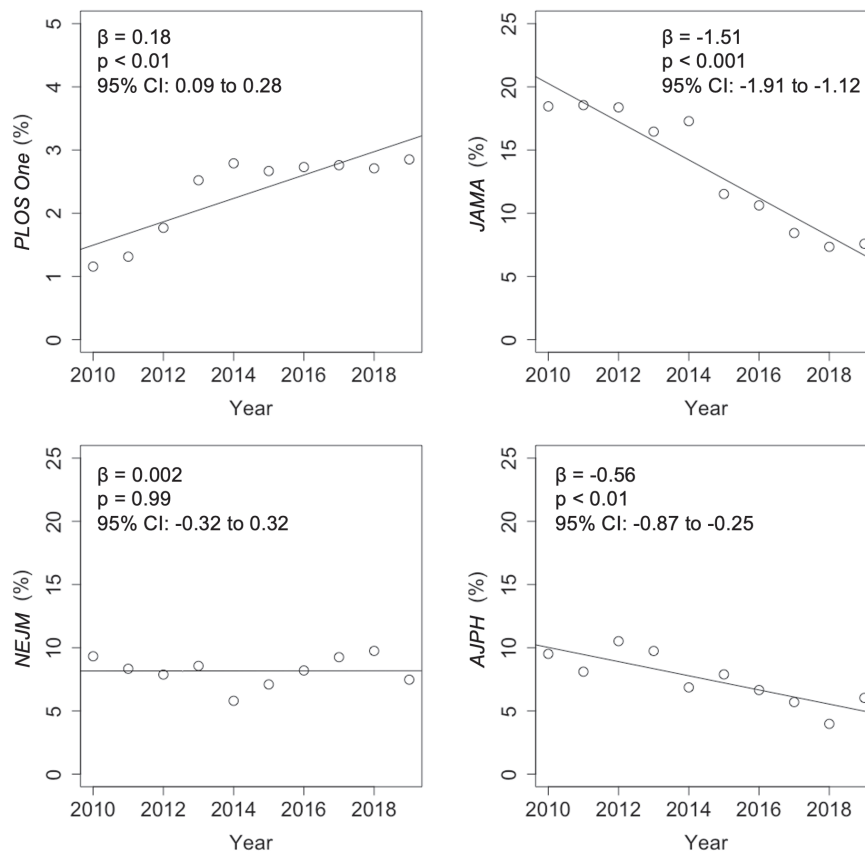
Subsequently, we evaluated whether the reported effect sizes were *correct* using criteria for the reporting of effect size from ORs developed from Davies *et al*<sup>6</sup> and Norton *et al*<sup>7</sup> (table 1). A *correct effect size interpretation* accurately reflects the definition of the ratio of odds. For example, if authors reported an OR of 1.5, the interpretation “50% increase in odds” is correct, while “50% more likely” is incorrect. Correct interpretations could also include other interpretations resulting from logistic regression (eg, change in probability, marginal risk).<sup>9 10</sup> Interpretations were independently reviewed by two authors (BC, ML;  $\kappa=0.76$ ). Disagreements were discussed with a third author (JWA, ECL) until consensus was reached. Articles with any incorrect reporting of effect size were labelled as incorrect, even if other ORs were interpreted correctly.

### Analysis

We computed the per cent of matching articles, including bootstrapped CIs, for the primary outcomes: (1) reporting the effect size from an OR and (2) providing a correct effect size interpretation. To evaluate differences in OR reporting across journals, we performed  $\chi^2$  tests. We evaluated if reporting practices were different among case-control studies (those described by authors as “case-control” or “case control” among our random sample) compared with all other studies using  $\chi^2$  tests. To quantify errors introduced by incorrectly reporting the effect size of ORs, we



**Figure 1** (Left) Trends in percentage of research articles that report the magnitude of effect size of an OR. (Right) Trends in percentage of research articles interpreting an OR that do so correctly. Data are summarised into 2-year intervals due to the low number of articles meeting the criteria.



**Figure 1** Trends in percentage of research articles that are searchable with keyword “odds ratio”, by journal.

calculated correct interpretations using methods and results reported in the article.<sup>8</sup> We used R 3.6.1 (R Foundation) for all analyses.

## Results

A total of 6124 articles with abstracts published in *AJPH*, *JAMA*, *NEJM* and *PLOS One* between 2010 and 2019 used ORs. At 13% (n=320), *JAMA* had the highest percentage of articles using ORs, followed by *AJPH* at 9% (n=361), *NEJM* at 8% (n=204) and *PLOS One* at 2% (n=5239). The proportion of articles using ORs in the past decade has declined in *JAMA* and *AJPH*, remained similar in *NEJM* and increased in *PLOS One* (figure 1).

Twenty-four per cent (94/400, 95% CI 20% to 28%) of sampled articles reported the effect size of an OR. Among articles making any effect size interpretation, 57% (54/94, 95% CI 47% to 67%) made an incorrect interpretation. Taken together, 10% (40/100, 95% CI 7% to 13%) of articles included a correct effect size interpretation of an OR. The proportion of articles reporting the effect size of an OR, and accuracy of such reporting has remained stable in the past decade (figure 2).

Examples of omitted, incorrect and correct interpretations are included in table 2. Most articles incorrectly reporting the effect size of an OR did so by misinterpreting ORs as risk ratios. For instance, one study found a result “8.3 times more likely” with an OR of 8.32, but when we accounted for the baseline prevalence of the non-exposed group (78.2%) the estimated risk ratio was 1.24. Another study interpreted that “risk ... is 100.7 times as high” based on a calculated OR of 100.7, yet we estimated the risk ratio to be 45.3.

Articles in *AJPH* more frequently ( $\chi^2=19.30$ ,  $p<0.001$ ) reported effect sizes (36%, 95% CI 27 to 45), than *NEJM* (26%, 95% CI 17.5 to 34.7), *PLOS One* (22%, 95% CI 13.9 to 30.2) and *JAMA* (10%, 95% CI 3.9 to 16.0). However, the probability of correctly presenting the effect size did not statistically differ by journal ( $\chi^2=0.56$ ,  $p=0.90$ ).

Articles reporting case-control design (n=42) were slightly less likely to report effect sizes (14%, 95% CI 4% to 24%) and had similar rates of reporting an incorrect effect size (50%, 95% CI 9% to 90%), compared with articles that did not report case-control design.

## Discussion

Articles in prominent journals often use ORs, but frequently omit reporting the effect size. Reported effect size interpretations were usually incorrect, in some cases by substantial margins.

Articles that omitted presenting the effect size typically reported only the associations implied by ORs and their statistical significance (eg, language such as “associated with”). Making informed interpretations thus falls onto readers, who may lack access to sufficient data to make actionable effect size interpretations. Many incorrect interpretations used language such as “probability” or “likely” when referring to the raw OR values, which could incorrectly portray ORs as risk ratios or likelihood differences. Interpreting ORs as risk ratios overstates effect sizes as events become more common; an OR of 2 could arise from event probabilities of 0.01 vs 0.005, 0.5 vs 0.33 or 0.8 vs 0.67, but the corresponding risk ratios would be 2, 1.5 and 1.2.<sup>7</sup>

Table 2 Example uses of ORs

Example quotes*	Effect size interpretation†	Correct interpretation‡	n (%; 95% CI)§
<p>“In adjusted analyses, gay and bisexual men were more likely than heterosexual men to have poor physical health (AOR = 1.38), disability (AOR = 1.26), and poor mental health (AOR = 1.77).”</p> <p>“Both 30-day mortality (OR, 0.76; 95% CI, 0.73 to 0.80) and 90-day mortality (OR, 0.73; 95% CI, 0.70 to 0.76) were significantly lower for azithromycin users.”</p> <p>“The primary outcome analysis showed a common odds ratio of improvement in the distribution of the modified Rankin scale score of 1.7 (95% confidence interval [CI], 1.05 to 2.8) favoring thrombectomy.”</p>	No	–	306 (76.5, 72.5 to 80.6)
<p>“This represented a 53% reduction in the risk of coronary heart disease among carriers of inactivating NPC1L1 mutations (odds ratio for disease among carriers, 0.47; 95% confidence interval [CI], 0.25 to 0.87; P=0.008).”</p> <p>“Households with a member who received information about chickens from a seminar are 8.3 times more likely to be aware of the vaccine (AOR: 8.32, 95% CI: 2 to 39, p&lt;0.01).”</p> <p>“Patients in the highest tertile of risk using the combined clinical and genetic model had a 7-fold increased risk of early stent thrombosis vs patients in the lowest tertile (OR, 7.63; 95% CI, 4.18 to 13.91).”</p>	Yes	No	54 (13.5, 10.1 to 16.9)
<p>“Each 1-SD increase in baseline log omega-3 fatty acid levels was associated with a 19% decrease in the odds of telomere shortening (unadjusted odds ratio, 0.81; 95% CI, 0.69 to 0.95).”</p> <p>“The marginal probabilities indicate that the percentage reporting fair or poor health was reduced by 4.8 percentage points (95% CI=0.8, 8.9; p=0.02) for current public housing residents.”</p> <p>“In contrast, we predict that a vaccination program would cause 41 excess hospitalizations (approximately 1 per 51 000 vaccinated infants) and 2 deaths due to intussusception in Mexico and 55 excess hospitalizations (approximately 1 per 68 000 vaccinated infants) and 3 deaths in Brazil.”</p>	Yes	Yes	40 (10, 7.0 to 13.0)

\*Each quote is an exact quote from an article meeting the inclusion criteria. Citations have been omitted to avoid singling out any given research group.

†Yes/No indicates if the study made an explicitly written interpretation of the magnitude of any reported OR.

‡Yes/No indicates if the study made a correct interpretation of the magnitude of any reported OR.

§Reports the corresponding n and prevalence with 95% CIs in parentheses.

AOR, adjusted OR.

Because our findings only included prominent journals and required only one interpretation of effect size, omissions and errors may be more common in the literature at-large. Indeed, misuse of ORs has been discussed in several specialty journals.<sup>11–13</sup> Such reports may have gone unheeded because they were construed as problems specific to subject specialties. Similarly, opinion pieces on the challenges of effect size reporting, despite publication in prominent journals,<sup>6,7</sup> may not have effected the necessary changes in practice without empirical data to support arguments. Our study is the first to systematically study how effect sizes are reported among studies in general health journals.

Our findings are limited by the use of PubMed to search for mentions of “odds ratios.” Because PubMed does not allow for searching through full texts of articles, randomly sampled articles may not represent all articles using ORs. While repositories like PubMed Central archive the full texts of articles, only open-access articles are indexed which excludes many recent articles in the selected journals. Furthermore, we only studied original research that included covariates, thereby excluding some articles with a single predictor variable, such as some randomised controlled trials. We limited our selection because studies with single predictors can report the effect size using descriptive statistics and use ORs to test statistical significance. In contrast, articles reporting original research with covariates must use the OR to report the effect size to control for confounding that exists within any bivariable comparison within the study. Lastly, to make our task feasible, we investigated only the use of ORs. Our findings highlight the need for additional investigation of effect size reporting for other statistics arising from binary, multinomial, count or continuous outcomes, as similar challenges may exist.

Omitted effect size reporting and erroneous interpretations may be due to the unintuitive nature of “odds”, which are more esoteric than *probability*.<sup>6</sup> Moreover, the magnitudes of ORs are incomparable across studies using different datasets and model specifications.<sup>7</sup> Specifically, ORs are conditional on the underlying

sample prevalence of the outcome *and predictor* (eg, changes in ORs may reflect changes in prevalence of predictors, rather than the underlying relationship with the outcome) and their relationship with covariates (eg, even when a variable independent of the outcome and predictor is added to the model, the OR changes).<sup>7,14</sup> In contrast, measures including *relative risk* (probability of outcome in exposed relative to unexposed) or *marginal effect* (difference in outcome, given change in predictor) are simpler to interpret and compare across studies thereby potentially leading to more frequent and accurate effect size reporting.<sup>8,10,14</sup> While certain study designs, such as case-control, must use ORs, the argument that reporting ORs alone sufficiently represent interpretable effect sizes is challenged by our findings of frequent erroneous interpretations, even in articles reporting case-control studies.

The consequences of not reporting or misinterpreting effect sizes may be compounded as results move beyond journals. Decision-makers (eg, legislative staff) and research disseminators (eg, news reporters) may lack training to describe the effect size from raw statistical results or to correct misinterpretations.<sup>15</sup> Policies and disseminations, once made, are often unobserved by researchers or editors and cannot be easily modified. This is particularly concerning for journals representing best practices in medicine (*NEJM*, *JAMA*) and public health (*AJPH*) that directly reach decision-makers and news reporters.

New standards to improve effect size reporting and its accuracy should be considered. For example, editors could require interpretations of effect size and evaluate their accuracy when evaluating submissions,<sup>3</sup> including requesting supplemental or alternative metrics facilitating ease of effect size interpretation like relative and absolute risks.<sup>8,10</sup> We urge developers of reporting guidelines to consider the importance of effect size interpretations, as current guidelines such as STROBE do not require them<sup>16</sup> and develop best practices for their use. Making health research more interpretable will make it more actionable for the benefit of public health.

**Acknowledgements** We thank Bryan E Dowd, PhD; Richard S Garfein, PhD, MPH; Theo N Kirkland, MD; Matthew L Maciejewski, PhD; John McGready, PhD; Edward C Norton, PhD; Davey Smith, MD, MAS; and Daniel Werb, PhD, for helpful feedback on earlier versions of our study.

**Contributors** BC, ML and JWA initiated the project and led the design. BC led the data collection, and all authors participated in the data analysis. All authors participated in the drafting of the manuscript, read and agreed to the final submission. BC, ML and JWA initiated the project and led the design. BC led the data collection, and all authors participated in the data analysis. All authors participated in the drafting of the manuscript, read and agreed to the final submission.

**Funding** The authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors.

**Competing interests** None declared.

**Patient consent for publication** Not required.

**Provenance and peer review** Not commissioned; externally peer reviewed.

**Data availability statement** The data used in the study are public in nature. The strategy to replicate our database is available in the text and the data are available on reasonable request from the authors.

**Open access** This is an open access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited, appropriate credit is given, any changes made indicated, and the use is non-commercial. See: <http://creativecommons.org/licenses/by-nc/4.0/>.

**ORCID iD**

Brian Chu <http://orcid.org/0000-0002-0803-5529>

## References

- Ioannidis JPA. Why most published research findings are false. *PLoS Med* 2005;2:e124.
- Gelman A, Stern H. The Difference Between “Significant” and “Not Significant” is not Itself Statistically Significant. *Am Stat* 2006;60:328–31.
- Harrington D, D’Agostino RB, Gatsonis C, *et al*. New Guidelines for Statistical Reporting in the *Journal. N Engl J Med* 2019;381:285–6.
- Chavalarias D, Wallach JD, Li AHT, *et al*. Evolution of Reporting *P* Values in the Biomedical Literature, 1990–2015. *JAMA* 2016;315:1141.
- Schwartz LM, Woloshin S, Dvorin EL, *et al*. Ratio measures in leading medical journals: structured review of accessibility of underlying absolute risks. *BMJ* 2006;333:1248.
- Davies HT, Crombie IK, Tavakoli M. When can odds ratios mislead? *BMJ* 1998;316:989–91.
- Norton EC, Dowd BE, Maciejewski ML. Odds Ratios—Current best practice and use. *JAMA* 2018;320:84.
- Zhang J, Yu KF. What’s the relative risk? A method of correcting the odds ratio in cohort studies of common outcomes. *JAMA* 1998;280:1690.
- King G, Tomz M, Wittenberg J. Making the most of statistical analyses: improving interpretation and presentation. *Am J Pol Sci* 2000;44:347.
- Norton EC, Dowd BE, Maciejewski ML. Marginal Effects—Quantifying the effect of changes in risk factors in logistic regression models. *JAMA* 2019;321:1304.
- Holcomb WL, Chaiworapongsa T, Luke DA, *et al*. An odd measure of risk: use and misuse of the odds ratio. *Obstet Gynecol* 2001;98:685–8.
- Tajeu GS, Sen B, Allison DB, *et al*. Misuse of odds ratios in obesity literature: an empirical analysis of published studies. *Obesity* 2012;20:1726–31.
- Katz KA. The (relative) risks of using odds ratios. *Arch Dermatol* 2006;142:761–4.
- Norton EC, Dowd BE. Log odds and the interpretation of Logit models. *Health Serv Res* 2018;53:859–78.
- Sutherland WJ, Spiegelhalter D, Burgman MA. Policy: twenty tips for interpreting scientific claims. *Nature* 2013;503:335–7.
- von Elm E, Altman DG, Egger M, *et al*. The strengthening the reporting of observational studies in epidemiology (STROBE) statement: guidelines for reporting observational studies. *PLoS Med* 2007;4:e296.