A New Inference about Adjuvants and Postoperative Pain Dosing?

##### Authors

Michael H. Andreae, M.D., Department of Anesthesiology & Perioperative Medicine,H187, Penn State Health Milton S. Hershey Medical Center, Penn State College of Medicine, Hershey. PA

Nathan L Pace, MD., MStat., Department of Anesthesiology, Univeristy of Utah, Salt Lake City, UT

##### Corresponding author:

Michael Andreae, MD, Address: Department of Anesthesiology, Department of Anesthesiology, Penn State Health Milton S. Hershey Medical Center, 500 University Drive, Hershey. PA 17033, Tel: +1 (717) 531 6140, Email: [mandreae@pennstatehealth.psu.edu](mailto:mandreae@pennstatehealth.psu.edu)

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# Editorial Text

Based on the best available evidence, which analgesic adjuvant is best for a particular patient to optimize postoperative pain control? For example, how much opioid sparing will the administration of intraoperative intravenous acetaminophen afford for a typical bariatric patient at Penn State Hershey or University of Utah Medical Centers? With patients and surgical procedures across the United States as diverse as species in the rain forest, this can be a difficult question. Often randomized controlled trial yield varying estimates of the clinical effectiveness of analgesic adjuvants. RCTs results, seemingly at random, leave clinicians baffled. Doleman et al in the study “Baseline morphine consumption…” propose an ingenious new solution to solve this riddle1. They synthesized the evidence for morphine dose reduction with adjuvants by controlling for baseline risk (morphine consumption) across surgical procedures and patient populations. With their novel approach, local audit data could be used to predict the expected average reduction in morphine consumption for any analgesic adjuvant. In the same breath, they modify and perhaps toss out the established paradigm of procedure-specific pain control.

Some variability in outcome estimates from RCTs investigating the same intervention in the similar population is to be expected. Each RCT recruits patients by a convenience sample; it is not a random sample of the entire population who might receive the adjuvant. By pure chance alone, each RCT has a chance to over- or to under estimate the effect of an intervention. Also, larger and smaller studies will lead to more or less precise estimates of effect of interventions. Meta-analysis pools available RCTs to synthesize the evidence and to increase the precision of effect estimates. This may reduce the uncertainty in the face of seemingly contradictory results. However, if the heterogeneity of the RCTs is very large (too inconsistent and disparate), then these summary statistics may be inappropriate and are frequently described as mixing apples and oranges. Excessive between-study heterogeneity in meta-analysis raises concerns that the included studies are clinically and methodologically too different, making pooling all identified RCTs unreasonable.

To explain why RCTs yield contradictory results, we can group studies by surgical intervention. Stratification by surgical intervention makes clinical sense because between-study heterogeneity may be smaller within each stratum. This led to the current paradigm of procedure-specific pain control2: It is expected that different surgical procedures cause different amounts of pain. Populations undergoing different interventions for different diseases may vary in how they respond to pain, in their comorbidities, pharmacokinetics etc. Clearly, a population of elderly men following thoracotomy for lung cancer will differ from a population of young women after cesarean section. It follows that postoperative pain control should be tailored to the specific surgical intervention and the particular population. While this seems intuitive, it drastically reduces the number of available studies for clinical decision making on for a particular patient population undergoing a particular procedure. For example, on thyroidectomy Doleman et al.1 found only one single RCT investigating the effect of intravenous acetaminophen. This poses a significant challenge to any evidence-based approach to procedure-specific postoperative pain management.

For studies investigating adjuvants for improved postoperative pain control, the effect estimate is the mean difference in morphine dose between those receiving and those not receiving the adjuvant. The mean dose of those not receiving the adjuvant can be considered to be the baseline risk for pain. Doleman et al.1 used Bayesian hierarchical meta-regression modeling adapted to requirements of such evidence synthesis3. They demonstrated that surgical procedure is not the main predictor of between-study variability in effect estimates for analgesic adjuvants to control postoperative pain, but baseline risk is. Restated, the benefit of the analgesic adjuvant is mainly driven by baseline morphine consumption in the control group and not by the surgical procedure. Two figures using acetaminophen data as an example will illustrate their findings.

Doleman et al.1 found 25 RCTs investigating intravenous acetaminophen for postoperative pain control. In Figure 1, a classical forest plot is shown, ordered by surgical interventions.[[1]](#footnote-2) The 25 RCTs have different effect sizes; the results are inconsistent, varying widely even within the same surgery.

In Figure 2, a schematic of the acetaminophen subplot of Doleman's Figure 21 is rendered.[[2]](#footnote-3) Each study is represented by a colored icon. If surgery were the best way to explain why studies yield different results, then studies investigating the same surgical procedure should have similar results. Hence the same color icons should be grouped together (along the same value on the y-axis indicating the estimated effect). But studies in the same surgical group are far apart on the y-axis. By contrast, studies with similar baseline risk (close on the x-axis, i.e. baseline morphine consumption in the control group) tend to have similar effects, (they are close on the y-axis, i.e. mean morphine-equivalent reduction), illustrated in a neat alignment along the regression line in Figure 2. The conclusion is that a meta-regression controlling for baseline morphine consumption in the control group is superior to stratification by surgery in explaining between study variance in results.

Doleman et al.1 propose that this is true for all analgesic adjuvants across all populations, for all surgical interventions, as shown in the other subplots of Doleman’s Figure 21. This contradicts the dominant paradigm of procedure-specific postoperative pain control. What is more, Doleman et al.1 also propose that clinicians can use local audit data, collected on local surgical populations, to estimate the average reduction of morphine consumption for each adjuvant. This is a great feature to have for a new unifying theory on how to synthesize the evidence for postoperative pain control.

Doleman et al.1 fit both a classical model and a Bayesian model that differs from the classical approach (also known as frequentist statistics). The Bayesian approach incorporates prior information and combines it with the newly observed data, much like a physician would in clinical practice. Both frequentist and Bayesian statistical models are based on subjective assumptions; the model choice itself is often more important for correct inferences. In both types of modeling, the analyst should explore the sensitivity of results and inferences to assumptions and model choices. In an available supplement, Doleman et al.1 provided software code, data and model details. They used very weak prior assumptions and in their sensitivity analysis found their results to be robust. They made their Bayesian approach transparent, sound and trustworthy.

Based on their previous work7, they speculated that baseline risk might better explain variability. Baseline risk for pain may modify the treatment effect of analgesic adjuvants as a proxy for unmeasured patient-level characteristics7. Their conjecture is supported by their meta analyses. This now suggests a set of testable hypotheses and predictions, making their proposed novel and unifying theory attractive. Progress in science and medicine is made by proposing models and hypotheses and then attempting to falsify these. Indeed, investigators can use local audit data to predict the reduction in morphine requirements and then prospectively observe the actual reduction achieved to investigate how well the Doleman model’s predictions hold. Meta-analysts can replicate the approach by Doleman et al.1 and for example test if it can be extended to evidence synthesis of regional anesthesia to explain and/or reduce the between study variability of the effects of regional anesthesia versus conventional multimodal pain control for postoperative pain control. If Doleman’s hypothesis holds, it would greatly simplify evidence synthesis and provide clinicians with much stronger and much more granular evidence on what analgesic to use for which patient. In fact, in the second part of their paper, Doleman et al.1 go on to use meta-regression to discuss rather refined and specific questions. For example, the question, if acetaminophen is more effective if administered intravenously rather than orally, is of great financial and clinical importance. But there are simply too few studies to adress this question, if we insisted on combining only studies comparing the effects in the same population/surgical intervention. Doleman et al.1 provide a league table of efficacy of analgesic adjuvants.

Doleman et al.1 acknowledge the many limitations of their work. Their meta-regression, like any meta-analysis was an ex post observational study, even if it had an a priori defined analysis plan. The confidence of their predictions and inferences will be weaker for ranges of baseline risk (baseline morphine consumption in the control group) where they have less studies for the particular analgesic adjuvant at hand. We will list these limitations pars pro toto using acetaminophen as an example:

* The range of surgical procedures included in the acetaminophen trials was limited, (see Doleman’s Table 11).
* The patient mix in the acetaminophen studies are predominantly female.
* The standard morphine dose (50 mg) at which dose reduction was estimated is much higher than the mean dose (28 mg) in the included acetaminophen studies. Inferences and predictions in the higher dose ranges therefore rely on fewer studies and/or on problematic extrapolation.
* For most adjuvants studied, there was little benefit (almost equal benefit) for studies reporting low baseline consumption, which is evidenced in Figure 2 of Doleman’s article1.
* Finally, the estimates in their regression equation apply to aggregate groups not individual patients.

We and the reader may remain somewhat skeptical and cautious about the approach and inferences suggested by Doleman et al.1 Provocative as the novel hypothesis by Doleman et al.1 may be, it would afford great utility. It also exemplifies the benefit and potential of novel Bayesian meta-regression approaches to support clinical decision making.

# Figures

## Figure 1

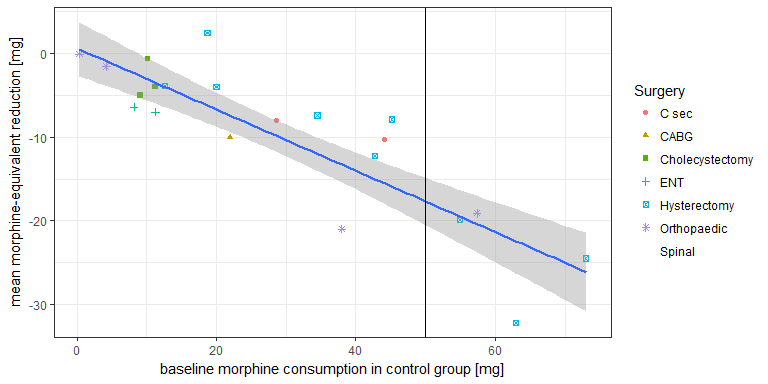


Figure 1

### Figure 1 Caption

Estimates for the mean reduction of morphine equivalent afforded by acetaminophen are shown in a classical forest plot with studies grouped by surgical interventions. The mean reduction of morphine equivalents and their 95% confidence intervals were provided by Doleman et al.. The later were used to estimate the standard error (Higgins 2011). The figure was produced with the Cochrane Collaboration software RevMan (RevMan 2014). The 25 RCTs have inconsistent effects for the drug under investigation (acetaminophen), varying widely even within the same surgery.

## Figure 2



### Figure 2 Caption

In Figure 2, we reproduce the acetaminophen subplot of Figure 2 of Doleman et al. Each study is represented by a dot. We coded and colored the studies by surgical procedure, according to the adjacent legend. If surgery were the best way to explain why studies yield different results, then studies investigating the same surgical procedure should have similar effects. Hence the same shape (and color) dots should be clumped together,(and the green-square-box cholecystectomy studies somewhat are). But most studies of the same color are far apart, for example orthopedic (violet-star) or hysterectomy (blue-cross-in-box) study dots are spread out over the entire range of the plot. By contrast, all studies line up neatly around the blue regression line, when we regress the mean reduction in morphine consumption (y-axis) against the baseline risk (baseline morphine consumption in the control group) in the x-axis. The conclusion is that a meta-regression controlling for baseline morphine consumption in the control group is superior to stratification by surgery in explaining between study variance in results.

# References

1. Doleman B, Sutton A, Sherwin M, Lund J, J W. Baseline morphine consumption may explain between-study heterogeneity in meta-analyses of adjuvant analgesics and improve precision and accuracy of effect estimates. Anesthesia and Analgesia 2017.

2. Kehlet H, Wilkinson RC, Fischer HBJ, Camu F, Prospect Working Group. PROSPECT: evidence-based, procedure-specific postoperative pain management. Best practice & research Clinical anaesthesiology 2007;21:149–59. Available at: <http://www.ncbi.nlm.nih.gov/pubmed/17489225>.

3. Sutton AJ, Abrams KR. Bayesian methods in meta-analysis and evidence synthesis. Statistical methods in medical research 2001;10:277–303. Available at: [http://journals.sagepub.com/doi/10.1177/096228020101000404 http://www.ncbi.nlm.nih.gov/pubmed/11491414](http://journals.sagepub.com/doi/10.1177/096228020101000404%20http://www.ncbi.nlm.nih.gov/pubmed/11491414).

4. Higgins JPT, Deek JJ. Chapter 7: Selecting studies and collecting data. In: Higgins JPT, Green S, eds. Cochrane handbook for systematic reviews of interventions 5.1.0 [updated march 2011]. Cochrane Collaboration, 2011:Available from www.cochrane–Available from handbook.org.

5. Review manager (RevMan) [Computer program]. Version 5.3. Copenhagen: The Nordic Cochrane Centre, The Cochrane Collaboration, 2014.

6. R Core Team. R: A language and environment for statistical computing. Vienna, Austria: R Foundation for Statistical Computing, 2013. Available at: <http://www.R-project.org>.

7. Achana FA, Cooper NJ, Dias S, Lu G, Rice SJC, Kendrick D, Sutton AJ. Extending methods for investigating the relationship between treatment effect and baseline risk from pairwise meta-analysis to network meta-analysis. Statistics in Medicine 2013;32:752–71. Available at: <http://doi.wiley.com/10.1002/sim.5539>.

1. The 95% confidence intervals provided by Doleman et al.1 were used to estimate the standard error.4 The figure was generated with the Cochrane Collaboration software RevMan5. [↑](#footnote-ref-2)
2. by the statistical software R6 with data provided by the authors.1 [↑](#footnote-ref-3)