## Where's the Beef?

How Much Can We Skimp on Pharmacokinetic—Pharmacodynamic Data?

Thomas K. Henthorn, M.D., Erik Olofsen, Ph.D.

Pharmacokinetic—pharmacodynamic models have been used extensively in clinical pharmacology to move beyond the doseresponse relationship to more directly relate drug concentrations to drug effects. <sup>1,2</sup> Pharmacokinetic—pharmacodynamic studies require plasma (or other relevant fluid) drug concentration measurements to characterize the pharmacokinetics as well as drug effect measurements, usually including onset and offset of the effect, to characterize the pharmacodynamics.

Use of pharmacokinetic-pharmacodynamic studies rather than dose-response studies is especially prominent in anesthesiology research, because anesthesia providers require the additional information about the time delay between attainment of plasma drug concentrations and their corresponding drug effects, and this

information is not obtainable with a mere dose–response study.<sup>3</sup> To fully capture this time delay (or hysteresis), pharmacokinetic–pharmacodynamic studies of anesthetic drugs require frequent plasma drug concentration measurements and drug effect measurements during both the onset and offset limbs of drug effect. This usually translates to obtaining both drug concentration and drug effect measurements during a drug infusion and for some time after discontinuing drug administration.

To conduct such a pharmacokinetic-pharmacodynamic study is resource-intensive given that it requires the timely collection of multiple blood samples for plasma drug concentration measurements, both of which are time consuming and expensive. In modern operating rooms, time is indeed money, and this maxim makes the conduct of



"We should be very cautious drawing conclusions in the language of pharmacokinetic pharmacodynamics when there are no drug concentration (pharmacokinetic) data." clinical studies in anesthesiology increasingly difficult even though the pharmacologic issues confronting contemporary anesthesia providers have become increasingly daunting.

A representative example of economized pharmacokinetic-pharmacodynamic studies appears in this issue of Anesthesiology in which the authors report on a clinical pharmacokinetic-pharmacodynamic study from the operating room environment addressing the clinically important question of whether patients with obstructive sleep apnea are more sensitive to the respiratory depressant effects of opioids, in this case remifentanil.4 If patients with obstructive sleep apnea are more sensitive to the effects of remifentanil, we would expect them to have a leftward shift in their remifentanil concentration versus minute ventilation relationship (or curve) com-

pared with controls. This leftward shift would show up as a statistically significant decrease in the concentration at the midpoint of the concentration-effect curve or the concentration producing 50% of the maximum effect (EC<sub>50</sub>).

The authors infused remifentanil at a constant rate for 10 min and measured minute ventilation *via* the anesthesia monitor at baseline and continuously during the 10 min of the remifentanil infusion, reporting the value at 5-s intervals. They economized on several key aspects of the usual pharmacokinetic–pharmacodynamic study paradigm. First, there were no plasma remifentanil concentration measurements. Instead, a published remifentanil pharmacokinetic model was used in place of an individual's remifentanil pharmacokinetics. Second, the clinical monitor reported a rolling 1-min average of minute ventilation rather than

Image: J. P. Rathmell.

Corresponding article on page 213.

Accepted for publication November 7, 2018. From the Department of Anesthesiology, University of Colorado School of Medicine, Aurora, CO (T.K.H.); Department of Pharmaceutical Sciences, Skaggs School of Pharmacy and Pharmaceutical Sciences, University of Colorado, Aurora, CO (T.K.H.); and Department of Anesthesiology, Leiden University Medical Center, Leiden, The Netherlands (E.O.).

Copyright © 2018, the American Society of Anesthesiologists, Inc. Wolters Kluwer Health, Inc. All Rights Reserved. Anesthesiology 2019; 130:186-9

snapshots in time or discrete time averages. Third, only the onset of the drug effect was measured because of the time constraints imposed by the operating room; the time course of the offset of effect was not evaluated. Fourth, although end tidal p $\mathrm{CO}_2$  measurements were obtained, hypothetical p $\mathrm{CO}_2$  values and their effect on minute ventilation were simulated, based on an indirect model of p $\mathrm{CO}_2$  and its effect on minute ventilation.<sup>5</sup>

Close examination of the raw data presented by Doufas *et al.*<sup>4</sup> indicates that their primary conclusion—that there was no significant difference between patients with and without obstructive sleep apnea in their respective sensitivity to the onset of ventilatory depressant effects of a 10-min remifentanil infusion—is correct. However, it is worth examining in detail the effects of some of the pharmacokinetic—pharmacodynamic study design restrictions<sup>6,7</sup> because the pressures to compromise on ideal study design are not likely to dissipate as further, increasingly difficult therapeutic dilemmas arise.

A pharmacokinetic-pharmacodynamic study has time and dose as independent variables and drug concentration and effect as dependent variables. If a study provides no drug concentration measurements, then drug concentrations are no longer a dependent variable and, instead, become an intervening variable, which is a hypothetical, unmeasured variable used to explain causal links between other variables, in this case between dose, time, and effect.8 Thus, changes in the magnitude and time course (kinetics) of the pharmacodynamic measurements from baseline will infer only the time course of changing drug input or drug concentrations because the latter are unmeasured. Verotta and Sheiner9 carefully examined this situation, pointing out the difficulties in identifying pharmacokinetic-pharmacodynamic model parameters in such circumstances. They proposed an inverse relationship between model or parameter identification and "the extent of knowledge of the input into the effect site." They proposed a hierarchy in which expense and information share a similar rank order: (1) experiments at steady-state conditions; (2) experiments in which concentrations are measured; and (3) experiments in which only the total input into the system are known. As Doufas et al.4 point out in their discussion, they have performed a type 3 experiment or a dose–response study.

Jacqmin *et al.*<sup>10</sup> considered type 3 pharmacokinetic—pharmacodynamic studies in which plasma drug concentration data are not available, and the principles they described have been used in the anesthesiology literature. <sup>11,12</sup> Jacqmin *et al.* termed this approach kinetic—pharmacodynamic as the "pharma," or drug concentrations, are absent. They relate drug infusion rate (rather than concentration) to a parameter they call EDK<sub>50</sub>. This is an effective dose or infusion (ED) that produces a half maximal response. The rate constant, K, indicates that there is kinetic variability in addition to the pharmacodynamic variability in the kinetics of drug

effect. Thus,  ${\rm EDK}_{\rm 50}$  is the product of the  ${\rm EC}_{\rm 50}$  and elimination clearance.

$$EC_{50} \cdot Cle = ED_{50} \cdot K = EDK_{50}$$

Although such an approach does not yield familiar terms such as EC<sub>50</sub>, it does retain the interindividual variability of both the pharmacokinetics and pharmacodynamics, even though their variability cannot be separated because of the inability to identify specific pharmacokinetic-pharmacodynamic model parameters in a type 3 dose-response study.<sup>9</sup>

Second, an issue with performing complex pharmaco-kinetic–pharmacodynamic experiments in a simple clinical setting is that the effect measurements afforded by clinical monitors may bias the estimates of model parameters. If a clinical monitor of minute ventilation reports a rolling 1-min average, then the values may have serially correlated residual error which, in turn, may affect estimates of interindividual variability. <sup>13,14</sup> Does this affect the conclusion of this study? No; but the extremely low estimate of interindividual variability in  $EC_{50}$  reported by Doufas *et al.* (7.9%) appears to be biased to a lower value by this mechanism and, therefore, should not be extrapolated beyond this study.

Third, a key aspect of meaningfully estimating the link between measures of drug input (e.g., drug concentrations) and drug effect (e.g., minute ventilation) in a type 2 experiment is closing the hysteresis loop of the drug concentration versus effect relationship. The differing concentration—effect relationships during onset and offset can be modeled, or mathematically reconciled, by  $\mathbf{k}_{c0}$  in a direct pharmacokinetic–pharmacodynamic model² or the combination of  $\mathbf{k}_{in}$  and  $\mathbf{k}_{out}$  in an indirect pharmacokinetic—pharmacodynamic model. Having concentration—effect data for both the onset and offset limbs of the hysteresis loop improves the confidence in the estimates of  $\mathbf{k}_{e0}$ ,  $\mathbf{k}_{out}$ ,  $\mathbf{EC}_{50}$ , and  $\mathbf{E}_{max}$ . Remember that  $\mathbf{EC}_{50}$  is the parameter estimate central to hypotheses that test drug sensitivity differences between groups in a type 2 experiment.

There are two modeling approaches to linking pharmacokinetics to pharmacodynamics. The method used most frequently in the anesthesiology literature is the direct approach in which a  $k_{\mbox{\tiny e0}}$  is a rate constant into and out of a hypothetical effect site, which closes the concentration-effect hysteresis loop, thus creating a hypothetical effect site concentration versus time curve. 1,2 The second approach is indirect pharmacokinetic-pharmacodynamic modeling, in which the observation of an effect (e.g., pCO<sub>2</sub>) is determined by the rate constants k<sub>in</sub> for effect production (e.g., CO2) and kout for effect elimination, respectively. 15 Both or either k<sub>in</sub> and k<sub>out</sub> can be affected by plasma drug concentrations. With an indirect model adjustment of  $k_{in}$  or  $k_{out}$  is sufficient to close the hysteresis loop, making a k<sub>e0</sub> and a theoretical effect site concentration redundant. To invoke remifentanil plasma and

effect-site concentrations Doufas *et al.*<sup>4</sup> used the direct pharmacokinetic–pharmacodynamic model of Minto *et al.*, <sup>16</sup> and to model the effect of pCO<sub>2</sub> on minute ventilation they used the indirect pharmacokinetic–pharmacodynamic model of Bouillon *et al.*<sup>5</sup> It is not clear what the effect on parameter estimation might be by combining direct and indirect modeling approaches.

It is possible to assess various modeling techniques by fitting simulated data to various models. To examine the issues introduced by scaled-down pharmacokinetic-pharmacodynamic study designs discussed earlier, existing population pharmacokinetic-pharmacodynamic models can be used to simulate remifentanil concentrations and remifentanil effects on minute ventilation to answer specific modeling questions. 17,18 We, therefore, generated data for 250 virtual patients, receiving a 10-min infusion of remifentanil, via Monte Carlo simulation with NONMEM 7.4.3 (Icon, USA) with random values generated using the pharmacokinetic-pharmacodynamic models of Doufas et al., 4 Minto et al., 16 and Bouillon et al. 19 The data for these 250 individuals were then fit to the following pharmacokinetic-pharmacodynamic models with (1) a fixed pharmacokinetic model (i.e., as Doufas et al.4 did), (2) an actual pharmacokinetic model (i.e., using each individual's pharmacokinetic model), or (3) no pharmacokinetic model (i.e., using a kinetic–pharmacodynamic model). As expected model selection criteria, using the -2 log likelihood objective function, showed that using a fixed literature-derived pharmacokinetic model is inferior and that there is a preference for the use of individual pharmacokinetic parameters based on actual, measured remifentanil concentrations. We found that the contribution of the interindividual pharmacokinetic variability to the estimate of the interindividual variability of the estimate of EC<sub>50</sub> had about the same order of magnitude as Doufas et al.4 reported.

The simulations also showed that there might be a preference for the simpler kinetic-pharmacodynamic model, as it was able to fit the data equally well. In the absence of plasma concentration data, as shown by Romberg et al., 12 a kinetic-pharmacodynamic model may capture the time course of effect measurements, although it cannot separate potential differences in pharmacokinetics from differences in pharmacodynamics. An interesting observation made by Doufas et al.4 was the lack of an inverse relationship between remifentanil effect site concentrations at the end of a 10-min infusion and minute ventilation (i.e., one would expect higher concentrations to be associated with lower minute ventilation; fig. 5a<sup>4</sup>). Our simulations verified the findings of Doufas et al. of a loss of correlation between the effect site remifentanil concentration at the end of the 10-min infusion when a fixed pharmacokinetic model is used. However, we found a clear relationship of decreasing minute ventilation with increasing remifentanil concentrations when simulated using each individual's pharmacokinetic parameters, and with the variabilities

set to those reported by Doufas *et al.*, but not when set to those of Bouillon *et al.*<sup>5</sup> Nonlinear mixed effects models, using NONMEM or similar software, have difficulty estimating data variability when data are correlated with each other.<sup>13</sup> Steps that account for serially correlated data may be required when devices such as clinical monitors that display rolling averages are used.<sup>14</sup> How rolling average (*i.e.*, correlated) data interplays with non–steady-state clinical studies and with fixed *versus* known pharmacokinetics deserves further study.

Near the dawn of pharmacokinetic-pharmacodynamic modeling, Verotta and Sheiner9 outlined the difficulties of describing the relationship between drug concentration and effect as investigators deviate from optimal study design. We are now entering an age in which there are libraries of population pharmacokinetic and pharmacokinetic-pharmacodynamic results, coupled with decreasing research funding and increasing clinical production pressures. The desire to marry literature-derived pharmacokinetics as an intervening variable with ordinary clinical data as dependent variables to address therapeutic dilemmas is understandable. But do we know enough about such reduced techniques to analyze these problems in the language of traditional pharmacology, using rate constants, Emays, Hill coefficients, effect site concentrations, and EC<sub>50</sub>s? Might deep "machine learning" or AI be more robust?20 Our opinion, based on the above analysis, is that we should be very cautious drawing conclusions in the language of pharmacokinetic-pharmacodynamics when there are no drug concentration (pharmacokinetic) data and when there is non-steady-state effect data and either the onset of effect or offset of effect is missing.

## Competing Interests

The authors are not supported by, nor maintain any financial interest in, any commercial activity that may be associated with the topic of this article.

## Correspondence

Address correspondence to Dr. Henthorn: thomas.henthorn@ucdenver.edu

## References

- Hull CJ, Van Beem HB, McLeod K, Sibbald A, Watson MJ: A pharmacodynamic model for pancuronium. Br J Anaesth 1978; 50:1113–23
- Stanski DR, Ham J, Miller RD, Sheiner LB: Pharmacokinetics and pharmacodynamics of d-tubocurarine during nitrous oxide-narcotic and halothane anesthesia in man. ANESTHESIOLOGY 1979; 51:235–41

- Sahinovic MM, Struys M, Absalom AR: Clinical pharmacokinetics and pharmacodynamics of propofol. Clin Pharmacokinet 2018; 57:1539–58
- Doufas AG, Shafer SL, Rashid NHA, Kushida CA, Capasso R: Non-steady state modeling of the ventilatory depressant effect of remifentanil in awake patients experiencing moderate-to-severe obstructive sleep apnea. Anesthesiology 2019; 130:213–26
- Bouillon T, Bruhn J, Radu-Radulescu L, Andresen C, Cohane C, Shafer SL: A model of the ventilatory depressant potency of remifentanil in the non-steady state. Anesthesiology 2003; 99:779–87
- 6. Paul M, Fisher DM: Pharmacodynamic modeling of muscle relaxants: effect of design issues on results. Anesthesiology 2002; 96:711–7
- 7. Avram MJ: Presenting data *versus* predictions as basic scientific information: Target-controlled infusions *versus* microgram per kilogram per minutes (reply). Anesthesiology 2011; 114:723; author reply 1–3
- 8. MacKinnon DP, Lockwood CM, Hoffman JM, West SG, Sheets V: A comparison of methods to test mediation and other intervening variable effects. Psychol Methods 2002; 7:83–104
- 9. Verotta D, Sheiner LB: A general conceptual model for non-steady state pharmacokinetic/pharmacodynamic data. J Pharmacokinet Biopharm 1995; 23:1–4
- Jacqmin P, Snoeck E, van Schaick EA, Gieschke R, Pillai P, Steimer JL, Girard P: Modelling response time profiles in the absence of drug concentrations: definition and performance evaluation of the K-PD model. J Pharmacokinet Pharmacodyn 2007; 34:57–85
- 11. Fisher DM, Wright PM: Are plasma concentration values necessary for pharmacodynamic modeling of muscle relaxants? Anesthesiology 1997; 86:567–75

- Romberg R, Olofsen E, Sarton E, Teppema L, Dahan A: Pharmacodynamic effect of morphine-6-glucuronide *versus* morphine on hypoxic and hypercapnic breathing in healthy volunteers. Anesthesiology 2003; 99:788–98
- 13. Karlsson MO, Beal SL, Sheiner LB: Three new residual error models for population PK/PD analyses. J Pharmacokinet Biopharm 1995; 23:651–72
- Nyberg J, Höglund R, Bergstrand M, Karlsson MO, Hooker AC: Serial correlation in optimal design for nonlinear mixed effects models. J Pharmacokinet Pharmacodyn 2012; 39:239–49
- 15. Jusko WJ, Ko HC: Physiologic indirect response models characterize diverse types of pharmacodynamic effects. Clin Pharmacol Ther 1994; 56:406–19
- 16. Minto CF, Schnider TW, Shafer SL: Pharmacokinetics and pharmacodynamics of remifentanil. II. Model application. Anesthesiology 1997; 86:24–33
- 17. Roozekrans M, van der Schrier R, Aarts L, Sarton E, van Velzen M, Niesters M, Dahan A, Olofsen E: Benefit *versus* severe side effects of opioid analgesia: Novel utility functions of probability of analgesia and respiratory depression. Anesthesiology 2018; 128:932–42
- 18. Henthorn TK, Mikulich-Gilbertson SK: μ-Opioid receptor agonists: Do they have utility in the treatment of acute pain? Anesthesiology 2018; 128:867–70
- Bouillon T, Schmidt C, Garstka G, Heimbach D, Stafforst D, Schwilden H, Hoeft A: Pharmacokineticpharmacodynamic modeling of the respiratory depressant effect of alfentanil. ANESTHESIOLOGY 1999; 91:144–55
- 20. Gambus P, Shafer SL: Artificial intelligence for everyone. Anesthesiology 2018; 128:431–3