

**Review essay on Rex B. Kline’s  
*Principles and Practice of Structural Equation Modeling*:<sup>1</sup>  
Encouraging a fifth edition**

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### Introduction

Kline’s fourth edition is reasonably strong but improvable. The text aims to introduce newcomers to fundamental structural equation modeling (SEM) principles, but tends to confuse “Principles” with “Rules.” Rules having insufficient grounding in principles leave readers ill-prepared for understanding and responding to changes in previously traditional “rules”—such as those concerning model testing, and latents having single indicators. SEM’s foundations would be clearer if Kline began by presenting structural equation models as striving to represent causal effects—a commitment that differentiates structural equation models from regression and encourages model testing. I begin this review by summarizing the covariance/correlation implications of three simple causal structures, which pinpoints multiple text improvements and underpins the discussions of measurement and model testing that follow. Causal structuring also grounds my later comments regarding modelling means/intercepts and interactions. A [file of Supplement Sections](#) expands on several points and lists multiple editorial corrections you might pencil into your copy of Kline’s text.

Kline’s fourth edition is more than one hundred pages longer than his third edition, and is effectively and compactly written. The material has been substantially reorganized, with the most substantive extension being a new chapter on “Graph theory and the structural causal model.” The publisher’s website contains syntax and output produced by an impressive variety of programs. Kline’s detailed discussion of several examples is noteworthy, and I count it as a strong positive that Kline’s examples include problems: “not all applications of SEM described in this book are picture perfect, but neither are actual research problems” (p. 1). We will encounter additional “problems” but if Kline prepares a fifth edition, I would recommend retaining the extra-problematic examples, along with supplemental discussions of what led to these slips, and instruction on ways to avoid similar slips. Overall, Kline’s text is solid enough to be worth improving. Regrettably, the tone of this review is more negative than I would prefer, but I could not find a way to detail the book’s positive features without also indicating some serious concerns. I asked Frank Trovato, editor of *Canadian Studies in Population*, to offer Rex Kline an opportunity to respond to my comments in hope that we might hear of Kline’s intentions regarding a fifth edition. I expect other readers of Kline’s fourth edition would appreciate your placing a reference to this review (and Kline’s response) in whatever copies you encounter. Indeed, my comments presume that you have access to the fourth edition for reference/comparison.

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1. Rex B. Kline. *Principles and Practice of Structural Equation Modeling*. New York: The Guilford Press, 2016. ISBN 978-1-4625-2334-4. Softcover US\$65, 534 pp.
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### The unavoidable implications of three simple causal models

If just variable  $X$  linearly causes  $Y$  with effect  $b$ , as in Figure 1A, this corresponds to the equation

$$Y = a + bX \quad (1)$$

and demands

$$\bar{Y} = a + b\bar{X} \quad (2)$$

$$\text{Var}(Y) = b^2 \text{Var}(X) \quad (3)$$

$$\text{Cov}(XY) = b \text{Var}(X). \quad (4)$$

The causal world makes variance in the causal variable  $X$  (namely,  $\text{Var}(X)$ ) produce, and thereby explain, variance in the effect ( $\text{Var}(Y)$ ). And the causal world makes variations in one variable ( $X$ ) produce coordination, correlation, or covariance ( $\text{Cov}(XY)$ ) between the causal variable and the effect. Variables' variances and covariances are consequences of causal actions, and we aspire to understand observed variances and covariances by locating the underlying causal structures. Observed covariances or correlations do not come from the math or statistics of equations; they come from the causal world that underwrites the equations. The causal world also coordinates the means of the variables (Equation 2). If the causal variable takes on a value corresponding to its mean ( $\bar{X}$ ), the resultant effect takes on a mean value ( $\bar{Y}$ ).

If  $Y$  has two correlated causes, as in Figure 1B, the relevant equation is

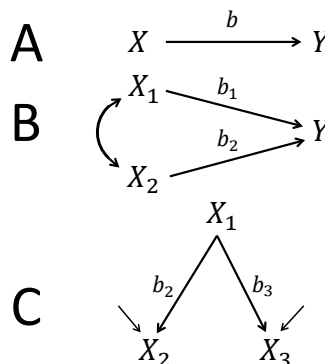
$$Y = a + b_1X_1 + b_2X_2 \quad (5)$$

and the causal world demands

$$\bar{Y} = a + b_1\bar{X}_1 + b_2\bar{X}_2 \quad (6)$$

$$\text{Var}(Y) = b_1^2 \text{Var}(X_1) + b_2^2 \text{Var}(X_2) + 2b_1b_2 \text{Cov}(X_1, X_2) \quad (7)$$

The partitioning of the causal world partitions the variance in the effect ( $Y$ ) but with the wrinkle that a portion of  $Y$ 's variance comes from coordination/covariance between the values of the causes, not merely from variations in the values of those causes. This variance equation is fundamental to: understanding why some explained variance cannot be uniquely attached to a specific cause, understanding how biased estimates result from omitting correlated causes, and understanding what goes awry if an error variable covaries with a cause (for example, if  $X_2$  was called a disturbance or error variable because it was not observed).



**Figure 1.** Basic causal structures.

Figure 1C introduces  $X_1$  as a common cause of  $X_2$  and  $X_3$ , where the causal equations are

$$X_2 = a_2 + b_2 X_1 + e_2 \quad (8)$$

$$X_3 = a_3 + b_3 X_1 + e_3 \quad (9)$$

With independent errors/disturbances, this causal structure has the unavoidable consequence that

$$\text{Cov}(X_2, X_3) = b_2 b_3 \text{Var}(X_1) \quad (10)$$

The  $X_2$  and  $X_3$  covariance is called *spurious* because no direct effect links  $X_2$  and  $X_3$ , but it is incorrect to describe the spurious correlation/covariance as “spurious (noncausal) associations” (p. 141). The association/covariance/correlation between  $X_2$  and  $X_3$  is the unavoidable consequence of the causal actions of the common cause, even if the relevant causal foundation is neither  $X_2$  nor  $X_3$  directly causing the other. The covariance in Equation 10 introduces the possibility of testing model implications, because estimates of the three right-hand terms can be obtained from the two model equations, and from data, without using  $\text{Cov}(X_2, X_3)$ . Comparing the model-implied covariance (from estimates of the two effects and the common cause’s variance) with the corresponding observed covariance between  $X_2$  and  $X_3$  might (or might not) challenge the depicted causal structure. The failure of specific model-demanded causal consequences to match with observed covariances underpins model testing and diagnostics striving to improve models’ causal structures.

The consequences of causal equations as presented above function the same way, whether the variables are observed or latent. And the consequences remain, even if some of the causally connected variables are observed while others are latent—which provides the principles grounding observed variables as measures of latent variables. Kline first introduces measurement of latents in Chapter 13 and in the context of factor analysis, where his emphasis on factors and outdated factor “rules” obscures the causal foundations of measurement, though measurement could have been helpfully introduced much earlier.

Equations 2 and 6 report that a case having a mean value on the applicable cause(s) is bestowed a mean value for the effect variable. Kline loses this easy and intuitive causal understanding when he turns to modeling means in Chapter 15, because he begins his discussion with non-causal regression. And he extends the confusion by referring to the  $a$  coefficients in these equations as “effects” of variables that are not variables “in the usual sense” (p. 371). That is, Kline omits cause from where it would be helpfully obvious, and adds cause where it really does not belong. Kline’s new Chapter 8 demonstrates an emerging acknowledgment of the relevance, utility, and unavoidability of causal understanding of structural equation models, but he has not yet incorporated that understanding consistently throughout his text. Beginning with a clear causal emphasis would strengthen the text’s foundational logic and encourage a focus on *principles* rather than *rules*. It would also reorient Kline’s discussion of model testing and diagnostics toward checking and improving the model’s postulated causal structures.

Readers seeking further instruction on the fundamental causal implications above, and unwilling to wait for Kline’s fifth edition, might see Hayduk (1987) Chapters 1 and 2, and Hayduk (1996) Chapters 1 and 2. All the variance and covariance equations for models structured as in Figure 1 can also be derived as special cases covered by the matrix Equation 4.30 in Hayduk (1987).

### Connecting the above to Kline’s text

Kline’s discussion of regression (Chapter 2) could have, and should have, differentiated between equations attempting to correctly represent a causal world and regression equations formed without requiring causal correspondence. SEM’s concern for proper causal specification is fun-

damental, and hence the associated pedagogical issues dig deep. Kline's Equation 1.2 (p. 13), for example, makes it seem like covariances are just statistical rearrangements of correlations, rather than being the consequences of causal forces as in Equation 10 above. Covariance does not come from correlation, as this equation seems to imply. Both the covariance and correlation are consequences of some underlying causal world, and the SEM researcher's task is to ferret out the nature of that underlying world. Understanding how underlying causal structures produce patterns in covariances is what makes it possible to "understand patterns of covariances" (p. 14). Similarly, Kline's Equation 2.2 (p. 26) seems to say that the structural coefficient on the left of the equation somehow comes from the correlation and other terms on the right, when the structural effect would in fact produce, and be the source of, the correlation. Kline's Equation 2.2 is not wrong, in the sense that the entities on the two sides of the equation really are equal, but the arrangement of the equation and its surrounding discussion obfuscate how causal action produces the correlation. Kline's Equation 2.3 is a rearrangement of Equation 2 for means above, but Kline's explanation—which is essentially an assertion that the equation holds, and his calling this a "mean structure" (p. 27)—provides no hint of how causal action links the variables' means, and similarly fails to ground the reader's understanding in the easy-intuition that a case having an average value on the cause should have an average value on the effect.

And consider whether a goal of structural equation modeling is to "explain as much... variance as possible" (p. 14), or whether the goal is to accurately determine how underlying worldly causal forces produce and hence explain variances, covariances, and means. Regression can be sold as attempting to explain as much variance as possible, but it would be preferable to present structural equations as focusing on how variance and covariance are explained. Focusing on how variance is explained would clarify that there are wrong ways and right ways of explaining variance. This, in turn, focuses attention on the correctness of the model's specification, and clearly differentiates SEM from regression by revealing *how* regression equations can be wrong as causal equations. This would make it possible to avoid multiple awkward transitions between what are supposedly "regression" equations and associated wordings that are expressly causal. Readers wishing to understand and monitor the multiple diverse consequences of Kline's failure to ground his text in a search for worldly causal structures should consider [Supplement Section 1](#). It is nice that readers report learning "something new" (p. 25) about regression from Kline's early material, but I would view it as more complimentary if readers had reported learning to differentiate between regression equations and structural equations pursuing causal understandings.

### Time sequence and causal action

In Chapters 6 and 7, Kline is inconsistent in his consideration of time and causal action. He claims "presumed causes *must* occur before presumed effects" (p. 123; emphasis added, and see p. 296, 432, 465). He also says, "the absence of temporal precedence may not always be a liability when estimating reciprocal causation" (p. 137) and includes examples of reciprocal effects (p. 135, 136, 143, 151, 152, 154, 156, 186). Causal actions are more easily recognized and estimated when the cause occurs first, but it remains possible to estimate reciprocal and looped causal effects (Rigdon 1995).

Kline's time ambivalence can also be seen in the awkwardness of his attempt to differentiate between mediation and indirect effects (p. 134), but it approximates contradiction when he claims that non-experimental designs "cannot establish which of two variables, a presumed cause and a presumed effect, occurred first" (p. 124–25). Valid reciprocal-effect estimates can be obtained *without* determining which occurred first (Rigdon 1995), and this undercuts Kline's idea that reciprocal effect estimates in non-experimental designs are "in some sense" "always wrong" (p. 137).

Indeed, SEM developed in the social sciences because SEM made it possible to investigate causal actions in non-experimental designs without confronting the ethical difficulties accompanying social experiments. Rather than claiming that effects operating at congruent times are inappropriate or impossible, Kline should have instructed his readers on model features that make it possible to estimate, and check, reciprocal or looped effects (Hayduk 1987, 1996; Rigdon 1995); he might even have discussed how causal loops provide reinterpretations of models previously estimated without loops (Hayduk 1996). Estimated models in which a variable directly causes itself (e.g., Hayduk 1985, 1996) demolish the supposed requirement of temporal precedence, because a variable can't precede itself! Count me *out* of the supposedly "emerging consensus that mediation analysis requires data from designs with time precedence" (p. 141; and see p. 465).

A related concern arises in when Kline argues that reciprocal causal connections between variables makes it "plausible that they may share unmeasured causes" (p. 138), and hence that the reciprocally connected variables should be assigned covarying disturbances. If  $Y_1$  causes  $Y_2$  with no disturbance covariance, there seems to be no general reason that adding a reciprocal effect from  $Y_2$  to  $Y_1$  should automatically manufacture the existence of a "new" common cause producing covariance between the variables' disturbances. New causal actions from a common cause (namely, a kind of causal structuring requiring a disturbance covariance) do not pop into existence merely because of the existence of some other causal action (even if that other causal action forms a loop or reciprocal effect). Similarly the "bows" in Figures 7.1a, 7.2, and 10.7, and on page 143, are *not* necessitated by the reciprocal or loop causes in the figures, and the disturbance covariance reported in the last line of Table 14.6 (p. 351) lacks justification.

### Observed variable and latent variable models

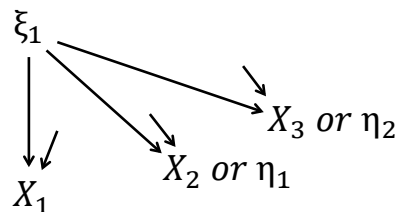
Chapters 6 and 7 consider the specification and identification of observed variable models, while Chapters 9 and 10 consider models containing latent variables. (The intervening Chapter 8 is addressed below.) These chapters are structured differently than in Kline's prior edition, and are afflicted by problems originating in: (a) the attempt to separate measurement from latent-to-latent effects (which obscures the advantages provided by modeling latent and observed variables simultaneously); and (b) the outdated presumption that latent variables require multiple indicators (p. 93). The chapter separation reflects the historical divide between path models and factor models, and seems to divide the rules for model identification into manageable chunks, applicable to first one part and then the other part of a model (p. 217). Unfortunately for Kline, the key advance provided by structural equation modelling was that it *overcame* the historical separation of measurement (via factor modelling) from structure (via path modelling) by combining and integrating measurement and structure. Some "strange" consequences of Kline's backsliding are presented in [Supplement Section 3](#).

Kline's rules for separately identifying path-like and factor-like model segments are insufficient for full structural equation models, and new "rules" will be required for overall model identification. Model identification rules have lagged behind the melding of factor and path model components, and lagged even further behind for models containing: fixed coefficients (e.g., fixed measurement error variances), constraints between coefficients, causal loops (whether longer loops or self-loops), means, intercepts, moderators, multi-level components, and latent variables having no direct indicators. It is reasonable to attempt to ensure coefficient identification, but Kline seems to employ his rules as instructions limiting how to build models (p. 119), rather than granting researchers' theory and hunches primary control.

Kline displays considerable inconsistency in how he specifies and applies his rules. Sometimes "a single indicator is preferred" (p. 217); meanwhile, his identification Rule 9.1 (p. 201) requires three indicators for a single latent factor, or two or more indicators for each of two or more correl-

ated factors. Thus, three indicators are identified, two indicators can be identified, and a single indicator may be preferred, yet we also read: “A better practical minimum is three to five indicators for each anticipated factor” (p. 195); “multiple-indicator measurement... is a cardinal characteristic of latent-variable models” (p. 127); and “each factor should have at least three indicators” (p. 454). If three indicators were really required, many of the models in Kline's Chapter 10 would be underidentified, because they contain latents having only two indicators—but they actually are identified, even though this is not evident because none of these two-indicator models were estimated.

Kline slants his advice to favour multiple indicators, but let's consider the second part of Kline's Rule 9.1 (p. 201)—namely, that a model would be identified with two or more correlated factors having two (or possibly more) indicators each. This identification “rule” really is *not* a general SEM rule, because SEM latent variables need not be “factors.” Kline connects his Rule 9.1 to CFA (confirmatory factor analysis), but how is a reader new to SEM supposed to understand that latents need not be factors, and that factor models lack the latent level effects and constraints that potentially make measurement errors on even single-indicated latent variables identified? Figure 2 illustrates how latent causal connections can assist measurement identification in much the same way as do additional indicators. The measurement error variance for a single indicator (like  $X_1$  in Figure 2) is often underidentified (unless provided a fixed value), but may be identified if the measured latent variable causes two or more latent variables, like  $\eta_1$  and  $\eta_2$ . If the two causally downstream latents in Figure 2 are well identified and do not influence one another, this causal structure mirrors the three-indicator identification condition reported in the first part of Kline's Rule 9.1. Focusing on the consequences of causal actions makes it easier to appreciate the parallel between downstream-latents and downstream-indicators, while avoiding causal action and emphasizing the difference between latent and observed variables obscures the parallel. Identification of SEM measurements is not just a matter of a latent and its direct indicators. Measurement identification relates to how the latent fits into a causal network composed of both its indicator(s) and other latents.



**Figure 2.** Latent effects potentially identify a single indicator's measurement error variance.

Failure to appreciate how latent level structure can assist in estimating and validating measurement persists into Chapter 16 on measurement invariance and multiple-samples, and robs Kline of an opportunity to free SEM from some of its historical factor analytic entanglements. Kline understands that structural equation modelling is moving away from traditional EFA and CFA (see his identification “Rules for Nonstandard CFA Models,” p. 202–06), but he seems not to recognize how the many other kinds of identification complexities render oblique rotations (p. 193), and rotational indeterminacy (p. 192) trivial tangents.

Concern for identification is relevant throughout the modelling process, not as in Kline's Figure 6.1 where identification is supposedly determinable *before* the selection of measures and remains unaltered by subsequent model revisions. Future texts would do better to present: the features that assist identification, the features that complicate identification, program output likely to appear for underidentified models, and ways of improving model identification (primarily introducing additional model or data constraints). I encourage readers to seek the structural model

features underlying Kline’s “rules” rather than memorizing the rules. Indeed, even models satisfying all the available identification rules can end up empirically underidentified due to random data variations or extreme values (p. 157, 206, 463), so even avid rule-followers should learn to recognize and respond to model underidentification.

A useful additional discussion of identification would address instances where the worldly model is underidentified given the researcher’s limited data. Important disciplinary issues arise if the researcher’s model is identified but the underlying worldly model is not identified given the available indicators. The more complex the causal world, the more likely it is that even nearly properly causally specified models will be underidentified, unless the researcher proactively addresses identification by employing causal variables entering the model at clear/precise locations, and constraining coefficients based on methodology or established “facts.”

Chapters 9 and 10 introduce latent variables—first as factor-based measurements and later as effects between latents. Kline would have served his readers better had he begun by considering measurement of latents without factors. This would have forced a consideration of the difference between a factor and a latent variable. A *latent variable* is a variable or characteristic whose true values are presumed to exist and hopefully impact some indicator, though other “error” variables’ causal actions prevent the latent’s values from fully determining the indicator’s values. Kline presents latent variables (Chapter 9, and p. 12–13) as if they are characteristics requiring multiple indicators—namely, as if they correspond to *factors*. Multiple indicators for a latent variable may be possible but are not required. Beginning with a latent variable like “age”—where true age differs from reported age (due to year-end jumps in reported age, memory, and avoidance of the next decade) would clarify that latent variables can have single indicators, and that researchers should address measurement error even with single indicators. Kline is simply wrong when he claims “multiple-indicator measurement...is a cardinal characteristic of latent-variable models” (p. 127; see also p. 213 last line, p. 220 second-last paragraph, and p. 223 third-last line). It is *not* the multiplicity of indicators that provides for latent variables. Latents are grounded in the acknowledgment of, and adjustment for, measurement error—where acknowledgment of, and adjustment for, measurement error can and should be done even with single indicators (Hayduk 1987, 1996; Hayduk and Littvay 2012). Latent variables are not necessarily factors, though *factors with multiple indicators* remain one style of latent variable. Second-order factors are latents having no direct indicators at all, and models may contain non-factor latents having no direct indicators (Hayduk 1990, 1996: Chapter 3).

Kline further confuses measurement when he equates latents with *constructs* and then says that constructs have different *facets*, as if latent variables also have facets (p. 127). A latent variable is a single, skinny dimension or number-line, and pretending that a single dimension has facets is nonsense. Speaking of facets and constructs demands causal connections between different latents, not reference to a single unidimensional latent.

Kline raises a different style of concern when he says, “Exogenous variable  $X_1$  is assumed to be measured without error, an assumption usually violated in practice” (p. 213; and see similar statements p. 132, 352). This statement acknowledges measurement error but sounds as if the common failure to appropriately adjust for measurement error can be excused merely because many people have done this! By the next page we read, “Because is specified as exogenous, it is assumed to have no measurement error” (p. 214). Unfortunately, being exogenous does not alter the existence of measurement error, and hence Kline has turned the common violation from the previous page into an “assumption,” as if this does not become a “usually violated” and deficient assumption. There simply is no justification for failure to compensate for a reasonable amount of measurement error in exogenous variables.

Measurement error in endogenous latent variables is equally important, though also miscommunicated by Kline. In reference to Figure 10.1a, Kline makes a statement that is appropriate for some endogenous variables (namely, those having no further downstream latent-level effects) as though it applies to all endogenous variables. “This assumption {namely of no measurement error} is not required for the endogenous variables in this model, but random error in  $Y_1$  or  $Y_3$  is manifested in their disturbances” (p. 213; { } material inserted). This statement is arguably true for  $Y_3$  in the relevant model, but it is definitely false for  $Y_1$ . For endogenous latents having no further latent effects (like  $Y_3$ ) the only consequence of measurement error variance is to increase the variance of the “disturbance” variable attached to the modelled endogenous variable (which makes the disturbance a mixture of measurement error and other omitted causes of the latent). The statement is false for  $Y_1$  in the Figure 10.1a model, because this variable’s measurement error functions *causally differently* (has different implications) than real omitted causes of the latent variable. Measurement error for a latent in the diagrammed  $Y_1$  position would causally impact only latent- $Y_1$ ’s indicator, while disturbance-style omitted causes of latent- $Y_1$  would impact both latent- $Y_1$ ’s indicator and the causally downstream variable  $Y_3$ . The differing implications introduce model misspecification, unless the model differentiates between measurement error and latent-level structural disturbance. (Kline makes the same mistake regarding  $X_1$  in Figure 10.2a, and  $Y_1$  in Figure 10.2b.) Rather than attempting to specify a new rule indicating when a researcher must adjust for measurement error in endogenous latent variables, it is simply safest to routinely acknowledge and adjust for measurement error in all latent variables all the time (Hayduk and Littvay 2012). And the adjustments should strive to attain validity, not merely “control for score reliability” (p. 127).

Contrary to Kline’s claim that “there is no point in retaining a model with just as many explanatory entities (factors) as there are entities to be explained (indicators)” (p. 190), there may be a very important point to be made. Differentiating causal actions dead-ending in each indicator (namely, measurement errors) from causal actions impacting the latent’s true values, which subsequently cascade to downstream latent variables, might require a model containing as many latents as indicators. The latents having single indicators would not be “factors” but SE models need not contain “factors.”

The diagrammatic partitioning of variance in Figures 6.3 and 9.2 (p. 131 and 190) differentiates between measurement error and real variations having unknown causes, but Kline pays insufficient attention to properly causally modelling the various variance components. For example, Kline recommends using scale reliabilities to determine fixed-measurement error variances (p. 223; and see p. 458). This adjusts for some measurement concerns, but is insufficient because proper modelling requires appropriate causal differentiation between all the indicators’ components (not just a reliability adjustment), and requires attention to their covariances, not just their variances. In the context of scales, the concern for validity recommends modelling all a scale’s items, not just scale scores.

### Factors versus factor scores, latents versus latents’ values

A different but related issue is whether SEM or SEM-measurements require that we know the values of latent variables. We sometimes would like to know the values of latents—for example, if the latent is someone’s ability and a related decision is required—but are we *required* to know the true values for latents in SE models? Think again of age. Are we required to determine the cases’ true ages in order to use latent age in a structural equation model? The answer is definitely no, we do not require the latent variables’ values. The complement of this is that we equally definitely do not need to know the values/magnitudes of error variables differentiating true latent variables’ values from observed indicators’ values. The statistical “magic” grounding the ability to estimate latent-level coefficients without knowing latent variables’ values can be seen in moving from Equation 4.28 to 4.29 in Hayduk (1987), but is a bit too cumbersome to detail here. The relevance of



this for Kline's text comes from *factor score* attempts to estimate the values of factor-latent variables. Factor score *indeterminacy* (p. 189, 193, 212) refers to the indeterminacy or imprecision in possible sets of estimated latent-factor values derived from the observed indicators. Kline says that "because theoretical variables and their proxies (indicators) are almost never identical, estimates of causal relations between latent variables are approximate at best" (p. 212). Let us repeat Kline's statement, replacing "theoretical variables" with "latent variables" and "proxies (indicators)" with "factor scores," estimated from the indicators, as Kline's context requires. "Because *latent* variables and their *factor scores* are almost never identical, estimates of causal relations between latent variables are approximate at best." To be even clearer, *because* we can't determine latent true scores precisely, estimates of the effects between latent variables are supposedly approximate at best.

Unfortunately, Kline's "because" is entirely unfounded. Obtaining proper effect estimates does not involve, require, or depend on latent variables' true scores. Contrast Kline's claim that latent-to-latent effects are "approximate at best" with the observation that latent effect estimates can be more accurate and precise because they adjust or compensate for measurement error, while the corresponding estimates from observed variables are prone to contamination with measurement error. (This was historically called *correcting for attenuation*.) Kline employs something that is not required in SEM (factor/latent scores) to inappropriately reframe a strength of SEM (adjustment for measurement error) into a supposed weakness!

Factor-score indeterminacy also seems to underlie Kline's warnings against "jingle-jangle fallacies" (p. 301, 458), where a single name is insufficient to force indicators to reflect a single latent (jingle), or different names are insufficient to force indicators to reflect different latents (jangle). More cogent warnings could have been made in the context of descriptions or interpretations of latents as encapsulating verbal understandings/theories, but where those understandings/theories may be inconsistent with the causal world controlling the indicators.

I end this section by noticing that factor-structured sets of indicators tend to fail because the required common-cause structuring is often inconsistent with the causal forces actually producing the indicator data. Kline's only real example of factor analysis appears at the end of Chapter 9 (p. 206–08) and is highly significantly inconsistent with the data. The inconsistency is not reported in the text, though it can be found in the publisher's model archive. Some sleight-of-the-writing-hand switched the strong *dis*-Confirmation of this factor model into a *Confirmation*, permitting the section to be titled a "CFA Research Example" (p. 206). Kline says that "relatively few applications of CFA are strictly confirmatory" because post-hoc model modifications are introduced to make the model fit, and/or because CFA is claimed after exploring with EFA, but it seems that even clear and direct model failure also fails to tarnish the *C* in CFA.

## Chapter 8, Structural causal models

It is a breath of fresh air to encounter Chapter 8's focus on causal actions and the unavoidable implications of causal actions. This chapter is new to Kline's fourth edition, presenting jargon introduced by Judea Pearl and "rules" reporting the unavoidable implications of directed acyclic graphs (DAGs). Kline makes a truly admirable attempt to present this material, but it is sufficiently complex that I offer some nit-picking suggestions (in [Supplement Section 4](#)), though I have two more substantial concerns. The first is that Kline does not introduce Pearl's *do(x)* operator (Pearl 2000: 70), which is fundamental to understanding the propagation of causal effects. The *do(x)* operator permits following the unavoidable consequences of *doing* or making specific, precisely expressed hypothetical interventions within the modeled causal structures. By excluding this, Kline loses an opportunity to assist his readers to see SEM's fundamental causal precision in action. The second concern is that Kline seems to have missed the consequences of Chapter 8 for his other chapters.

This causally focused chapter contrasts with Kline's repeated avoidance of cause, as discussed above. The precise and unavoidable causal implications discussed in Chapter 8 should have fortified Kline's Chapters 11 and 12 on model testing. Kline has argued against coefficient testing in other contexts (Kline 2013), but Chapter 8 provided a missed opportunity to clarify the fundamental difference between coefficient tests and model tests. DAG tests are tests of precisely demanded model/theory implications—namely, they are tests that incorporate and investigate the combined consequences of multiple theorized causal structures, and differ from direct tests of specific coefficients.

### Chapter 11, Estimation and local fit testing

Chapter 11 reads smoothly but is sprinkled with multiple problematic statements. One easily misconstrued claim is that single-equation estimation methods “can be less affected by specification error than simultaneous methods” (p. 231; and see p. 235). The misconstrual comes from failing to consider the other, not-less-affected equations, and from the single equation methods discussed being routinely misspecified because they fail to compensate for measurement error (see p. 233, point 1). There are single-equation estimation methods that address measurement (e.g., Bollen et al. 2014) but these are not discussed, and have their own limitations. Another misconstrual is embedded in Kline's description of maximum likelihood estimates as “the set of parameters estimates that is most likely to have generated the observed data” (p. 236). The misconstrual should become clear if you hear this as an example of “the false belief that  $p$  measures the likelihood that  $H_0$  is true, given the data” (Kline 2013: 98).

But a more fundamental concern is Kline's inappropriate adoption of a specific correlation-discrepancy size of .10 as his implicit *local fit test* criteria. Smaller discrepancies between observed and model-implied correlations are presumably acceptable, while discrepancies “of .10 or more may signal appreciable model-data disagreement” (p. 240). Kline repeatedly appeals to whether or not .10 is exceeded (p. 253–54, 278, 329, 380, 385, 408, 416, 481) and speaks as if this constitutes reasonable local fit testing (p. 241, 283). Unfortunately, exceeding or not-exceeding a correlation discrepancy of .10 does not constitute a statistical test, and the .10 value lacks statistical justification. The value merely demarcates the boundary between what Kline will attend to or disregard. (Actually, Kline does not even stick consistently to his .10 value (p. 344).) Programs like LISREL and EQS report standardized residuals (p. 252), which provide statistically appropriate local fit tests, but Kline displaces the available tests with a “criterion” more to his liking, even though he knows important model misspecifications can produce only smaller amounts of model-data discrepancy (he cites Hayduk 2014a), and even though he has heard that “Shame for disrespecting evidence {will constitute one of} the personal consequences of insufficient respect for structural equation model testing” (p. 496; {} material added).

Combining “Estimation and local fit testing” into a single chapter has the unfortunate consequence of placing local fit testing prior to overall model testing (namely, Kline's next chapter). Both model testing and local fit testing depend on estimation, but overall model testing should precede investigation of local fit. For a bad example, notice that after dedicating more than a dozen pages to a detailed example, Kline says “the fit of the example model is unacceptable” (p. 253) both locally and globally, but he fails to respond appropriately to the model-data inconsistency. Changing the model's structure to conform to the world's structure would: alter the control variables, alter the estimates, alter the estimate's significance, alter the basis sets, alter the residual ill fit, and mess with just about every claim Kline made about his example. Kline's comment regarding model-data inconsistency does not even hint at the numerous diagnostic investigations that should be undertaken, or the substantial model reassessments that should accompany detection of model-data inconsistency. It is easy—deceptively easy—to think that if a model “poorly

explains certain observed associations” (p. 253), the model problems are tightly linked to those particular problematic covariances/correlations rather than being detectable symptoms of more dispersed yet fundamental model misrepresentations. Localized ill fit does not confidently report localized model specification problems. Patterns of local ill fit can sometimes contribute usefully to diagnostic examinations, but even patterns of local ill fit do not irrefutably detect the relevant model problems. I count it as a serious deficiency that the term “diagnostics” does not even appear in Kline’s subject index. Diagnostics require assessing the many things potentially wrong with a model, and/or the data, rather than routinely freeing the nearest error covariance—which often amounts to blaming/convicting the nearest bystander.

### Chapter 12, Global fit testing

Chapter 12 also reads smoothly, but it is perhaps Kline’s most problematic chapter. The problems begin with the title. Kline knows “that there is actually no dependable or trustworthy connection between the size of the residuals and the type or degree of model misspecification” (p. 278)—where residuals refer to the difference between the model-implied covariances/correlations and the data covariances/correlations. Given the *disconnect* between the amount of ill fit and the seriousness of model misspecification, researchers face a choice of being interested primarily in model misspecification or model fit. Even a brief consideration determines the primary concern is model misspecification, while fit plays only a supporting role. Researchers want to test their *model*, not just their model’s fit, and examine fit to see whether or not this provides evidence of model misspecification. Thus, a more appropriate chapter title would be “Testing for model misspecification” or “Detecting model misspecification” rather than the current “Global fit testing.”

If ill fit is detected, researchers should probe the program output for diagnostic clues to the nature of the problem(s) and potential model or data improvements. A ringing ill-fit alarm bell should prompt thorough and detailed investigation of possible data mistakes as well as the multiple kinds of possible model misspecifications—not mere pursuit of different fit-index ways of reporting the ill fit (p. 266).

Kline grounds his fit-index-based disrespect for test evidence in an oft-cited claim by Box (1976: 792) that “all models are wrong” (p. 263)—which implicitly and inappropriately suggests that model misspecification cannot be avoided, so you shouldn’t worry if you encounter some. God might know whether or not all models are wrong, but how could even a famous person like Box know about all structural equation models—including models that have not yet been specified or run? In fact, Box was not even referring to structural equation models—he was writing back in 1976, when structural equation modelling was relatively unknown, and SE model testing nearly nonexistent. Incompleteness can make many styles of statistical models wrong, but incompleteness does not necessarily contribute to SEM ill fit, because measurement errors and unknown latent disturbances are *parts* of structural equation models, not omitted features. And given that we can construct structural equation models of experiments, applying Box’s statement to SEM implicitly asserts that all experiments are wrong—because SE modeling of experiments would also only result in wrong models! Claiming that all experiments are wrong is clearly nonsense.

None of these kinds of considerations have stopped Kline (and some others) from propagating this nonsense and its paraphrases. See: “When (not if)” the model does not fit (p. 120), models “are imperfect approximations” (p. 262), and fit indices “allow for an ‘acceptable’ amount of departure from **exact (perfect) fit**” (p. 60; emphasis in the original). Correctly specified models may be rare (p. 232), but rare does not mean impossible (for fitting models, see Entwisle et al. 1982; Hayduk 1994; Hayduk et al. 1997, 2005).

Kline ends his quote from Box (p. 263) a bit too soon. Box's next sentence reads: "Since all models are wrong, the scientist must be **alert to what is importantly wrong**" (Box 1976: 792; emphasis added). Kline stumbles regarding what is "importantly wrong" because he frequently conflates importance with the amount of ill fit, rather than with the nature of underlying model misspecifications. For example, Kline provides a section on a "Recommended Approach to *Fit Evaluation*" (p. 268; emphasis added) rather than to *Model* evaluation. By focusing on *Approximate Fit Indexes* (p. 266–68), RMSEA (p. 273–75), CFI (p. 276), and SRMR (p. 277), and continuing with these indices in his examples, Kline distracts from a search for "what is importantly wrong." Kline notes that there are "discredited thresholds for such fit statistics" (p. 269, 268), without reporting which specific thresholds have been discredited, and despite his continuing use of thresholds courting discreditation (p. 267, 274, 276–78).

Kline fails to appreciate the depth of the challenge to ALL model fit indices created by there being "no dependable or trustworthy connection between the size of the residuals and the type or degree of model misspecification" (p. 278). He proceeds as if small-sized residuals overrule or overturn the significance of those residuals, whether in the context of global fit testing or in local fit not-real-testing, via his indefensible .10 correlation residual (p. 462, 265). This can be seen in Kline's claim that a large  $N$  devalues  $\chi^2$  testing because a larger  $N$  enables  $\chi^2$  to detect smaller covariance/correlation residuals—including residuals smaller than his .10. For many *misspecified* models,  $\chi^2$  power does increase with  $N$  and thereby demonstrates increasing power to detect misspecifications, but for properly specified models,  $\chi^2$  does *not* increase with increasing  $N$  (Hayduk 2014b). Kline inappropriately reports  $\chi^2$  as being "overly sensitive to sample size" (p. 271; see also p. 330, 462), when in fact  $\chi^2$  increases with  $N$  only when there is some detectable problem in the model or data. Kline's Exercise 4 (p. 279, 298) and its suggested answer (p. 480) are ill-founded, because with a proper model and larger  $N$ , the data covariances would become more stable due to smaller sampling variations from the true covariances, and consequently  $\chi^2$  would not inflate. The idea that  $\chi^2$  is "overly sensitive" implicitly appeals to there being discrepancies that are too small to be worth detecting and investigating, when in fact covariance discrepancies can be zero even in the presence of important model misspecifications (Hayduk 2014a). Kline's proclivity to think of large- $N$   $\chi^2$  as detecting trivial fit differences parallels his tendency to disregard local ill fit correlations less than .10, even though there is no justification for claiming that smaller residuals in either context protect researchers from important model misspecifications.

A related imprecision is Kline's failure to distinguish between the causal structure of a model and the fit provided by that model. For example, Kline titles one section "Equivalent CFA models" (p. 315) and another "Equivalent SR models" (meaning *Structural Regression models*; p. 348), when what he is discussing are causally *non-equivalent* models providing equivalent fit. The models are not causally equivalent because they contain different causal specifications, though they have corresponding covariance implications. Had Kline's titles been something like "Different CFA or SR models producing equivalent fit," the discussion would have turned to causal specification/misspecification, and the models reported on pages 345 and 358 would have been described as *not distinguishable on the basis of their covariance fit*, even though the worldly causal structures are empirically distinguishable.

Kline's discussion of RMSEA (Root Mean Square Error of Approximation) contains multiple technically correct statements but is unlikely to assist anyone not already familiar with the RMSEA (p. 273), and actually encourages problematic SEM practice. Kline cites a 2008 work as indicating there is "little support for a universal threshold of .05 (or any other value)" for the RMSEA (p. 274), yet spends the next pages propagating obsolete threshold values suggested by Browne and Cudeck back in 1993, only to follow this with additional relatively recent references that "question the generality of thresholds for the RMSEA" (p. 276). As if these were not enough, Kline disregards the logical problem at the heart of the RMSEA.

The problem is that non-zero RMSEA criteria attempt to excuse or overlook some amount of real model-data inconsistency for each model degree of freedom. The RMSEA is calculated as *ill fit per degree of freedom*, and hence claiming a non-zero RMSEA value as acceptable claims that some non-zero amount of real ill fit is acceptable for each and every model degree of freedom. (The Browne and Cudeck reference Kline cites as foundational for the RMSEA describes real (non-random) model-data inconsistency as “error of approximation” (1993: 141).) Overlooking, excusing, or discounting real model-data inconsistency is clearly problematic. And a model rendered strongly testable by having many degrees of freedom is supposedly excused (ahem) from that strong-testing because Browne and Cudeck said that the RMSEA permits (cough) overlooking some amount of model-data inconsistency for each degree of freedom—an amount not based on statistics but “based on subjective judgment” (1993: 144).

Kline (p. 274) cites work documenting a clear instance where the same Michael Browne (of Browne and Cudeck), along with Robert MacCallum (another big name who championed disregard of real model-data inconsistency) and Kim, Andersen, and Glaser (2002), defended and retained a model that was inconsistent with their data. Browne et al.’s “supposedly negligible ill fit obscured important, systematic, and substantial causal misspecifications” that were located and corrected by attending to relevant experimental conditions (Hayduk et al. 2005: 1). Somehow Kline remains immune to the methodological unacceptability of overlooking real model-data inconsistency, despite: the “critical” RMSEA value being subjective (not statistically based), challenged by multiple recent references, and having led strong people into making indefensible modelling mistakes.

Unfortunately, Kline perpetuates this problem throughout the remainder of his book. It is nice that Kline’s testing chapter summary reports there being a “consensus that some routine practices are inadequate” (p. 297), and nice that his best practices chapter says, “Do not rely on ‘golden rules’ for approximate fit indexes to justify the retention of the model” (p. 461), but these come across as disingenuous, given that he also says, “If possible, report...the RMSEA” (p. 464), given that the logical problems with indices are not respected in Chapter 12, and given that the RMSEA is repeatedly incorporated later (p. 279, 290, 305–07, 317, 328, 344, 347, 350, 357, 359, 380, 385, 406, 416). The same inconsistency appears when Kline complains that low power provides “little chance of detecting a false model” (p. 265), while encouraging low power—by appealing to fit indices rather than tests, by denigrating the power provided by larger  $N$ , and by switching from a test of model fit to the hypotheses of “close fit” and “not-close fit” (p. 290–91). If he really appreciated power, he would have considered power in the context of the most powerful available test—namely, the  $\chi^2$  model test—not merely any lower-power index. In fact, power is defined only in the context of model testing, not indexing, so a statistical sleight-of-hand accompanies discussing the power to detect arbitrarily hypothesized index values (p. 292, Figure 12.2 and fourth line). In short, Kline has not yet come around to consistently reporting the deficiencies of the RMSEA and warning against its continued use.

Unfortunately, Kline’s text falters in several additional places due to deficient model testing, as you can see by considering the technical-teasers in [Supplement Section 5](#). I assisted the third edition of Kline’s *Principles and Practice of Structural Equation Modeling* by providing some “backbone” (Kline 2011: *xi*) to his testing chapter, only to find that his fourth edition reverted to the **deficient** view that there should be “LESS EMPHASIS ON SIGNIFICANCE TESTING” and that the “proper role for significance testing in SEM is *much* smaller” (p. 17; both emphases in the original). I hope the above constitutes a sufficiently clear and strong prosthetic to propel Kline, or his successors, into understanding the important difference between a model being significantly inconsistent with the data versus “acceptably close” to the data (p. 11).

## Chapters 13 and 14

“Confirmatory factor” and “Structural regression” models are addressed in Chapters 13 and 14, respectively. Factor analytic measurement of latent variables developed quite independently of regression/path-analytic linking of different variables, and this led to a common presumption that structural equation modelling should proceed in two separate steps—initial measurement of latent variables via factor analysis, followed by structural connections between different latents. Kline cites Anderson and Gerbing (1988) on two-step modelling and aligns himself with the two-step approach by separating these chapters. This supposedly reduces the complexity of model assessment by providing a “separation of measurement issues from structural issues” (p. 340).

Unfortunately for Kline, one of SEM's greatest strengths is to combine, not separate measurement and structure. The reason is simple. Structural equation modeling strives for *valid* models and *valid* measurement, not merely reliable models and reliable measurement. Measurement validity requires that a measured *latent variable function appropriately* in connection to *other latent variables*. The possibility of demonstrating appropriate connections to diverse theory-specified variables only arrives with the latent level of the model, and hence measurement remains incomplete until the measures are incorporated into full, well-functioning models. All of a latent's indicators (not scales or parcels) should be retained in the full SE model if a latent proposed by factor analysis seeks validation.

Kline's emphasis on multiple factor-structured indicators results in insufficient attention to measurement error variance in routinely used variables like age, sex, and education, which rarely have more than a single indicator. Another consequence of requiring multiple indicators is that this tends to displace latents which could function as control variables, instrumental variables, or variables clarifying the operative causal mechanisms. Kline cites Hayduk (1996) but somehow misses its Chapter 2, which explicitly challenges Anderson and Gerbing's two-steps; he also cites Hayduk and Glaser (2000a), but somehow also misses the corresponding challenges to four-steps. Kline is seriously off-base when he cites Hayduk and Glaser (2000a, b) as supporting either two- or four-step modelling (p. 339). In short, by recommending use of “two-step modeling, not one-step modeling” (p. 462), Kline renders his book incapable of providing a thorough discussion of measurement validity, burdens his reader with identification “rules” that really aren't sufficient-rules, misses adjustment for measurement error in variables like age, and hinders pursuit of informative model specifications.

In Chapter 14 we re-encounter the problem of waffling between causal and non-causal connections between variables. A conscientious reader will cringe at the chapter title's reference to *Structural Regression Models*, because structural effects at the latent level are not mere regression coefficients. Kline's sporadic attention to latent-level causal structuring is evident in his “Detailed Example” (p. 341–48), which considers a model by Houghton and Jinkerson (2007). Kline says these “authors describe the theoretical rationale for each and every direct effect among the four factors in the structural model” (p. 220), which exudes careful causal attention. Then we notice that four of the six latent-level effects in the model Kline “would retain” (p. 347) are insignificant (Figure 14.2, p. 348). The insignificant effects disconnect the *Constructive Thinking* latent from the remainder of the model's latents, including the final dependent variable *Job Satisfaction*. The absence of evidence supporting these effects contrasts sharply with Houghton and Jinkerson's claim that their study results “suggest that *constructive thought* strategies are related to *job satisfaction*” (2007: 51; emphasis added). Surely either the theory or measurement are problematic if the measurements do not locate latent variables displaying causal actions required by theory! Several modelled indicators had been constructed by segmenting sets of items into parcels, or sub-scales, whose reliabilities were previously reported (see p. 220; and Houghton and Jinkerson 2007) and several of the items had uncomfortably low proportions of explained variance. But the causal disconnection seems not to

have prodded any reconsideration of the measures. Somehow both the measurement and theory emerge unscathed, despite the vanishing theorized effects!

Further causal imprecision appears in this example when Kline adds a measurement error covariance between the indicators of Happy and Mood<sub>2</sub> because there is “common item content across the two indicators” (p. 344). Unfortunately, the estimated covariance is inconsistent with this common content theory, because it is negative (p. 347)! Common content would be consistent with a positive covariance, but not a negative, measurement error covariance. But why might Kline have missed this? Notice that Kline’s justification appeals to “common item content,” not to a common *cause* of the items. Had Kline thought of his justification as requiring a common cause, that should have triggered considerations paralleling Figure 1C and Equation 10 above, which report that a common cause only produces a negative covariance if the effects have opposite signs—but opposite signed effects are incompatible with Kline’s appeal to common content. The unjustified negative coefficient seems to “‘clean up’ local fit problems” (p. 345), and provides a clear example of how causally inappropriate and likely misspecified coefficients can deceive researchers by sopping up local ill fit. Sure, the fit is improved, but the properness of the model’s causal specification has been sacrificed to attain the improved fit.

Kline’s example contains yet another missed opportunity. The measurement error covariance that Kline added to his measurement model results in his model having one fewer degree of freedom than Houghton and Jinkerson’s corresponding model (Kline’s 47 versus their 48). Their final model restricts two additional effects and leads to Kline’s model having two more degrees of freedom (49); but Houghton and Jinkerson (2007: Table 2) report three more degrees of freedom (51 rather than 50) for their final model. Kline’s Chapter 6 exercises instruct readers in counting degrees of freedom, but Kline misses this opportunity to apply this skill. One latent variable in Houghton and Jinkerson’s final model seems to have been scaled in two different ways, namely, by setting a “loading” to 1.0 and by simultaneously setting the latent variable’s variance to 1.0, like all their other latent variables (Houghton and Jinkerson 2007: 47, Figure 1). This double-scaling forces their *Job Satisfaction* latent to contribute exactly 1.0 unit to the variance of their *Job Satisfaction* indicator (like setting both the variance and effect in Equation 3 above to 1.0 if *Y* was the indicator), when that indicator actually has a variance of  $0.939^2 = .88$ . This forces a model-data inconsistency, explaining why Kline’s model fits better than their model, and should have produced an impossible negative error variance estimate for Houghton and Jinkerson’s scaling indicator. Kline missed this opportunity to show how model misspecification can lead to problematic estimates, and missed the opportunity to caution that the literature contains enough errors to warrant routine caution and checking.

## Chapter 15

This chapter addresses modelling means and latent growth curves. It reads smoothly but requires revision because it both omits causation where it is appropriate, and inserts causation where it is inappropriate! Equations 2 and 6 above illustrate how intercepts coordinate the means of causally connected variables, and this parallels Kline’s Equation 2.1 (p. 369 and 27), but Kline flounders because his equation is presented in the context of regression rather than causal action.

In the example spanning pages 369–72, Kline interprets the intercept of 20.000 in his regression equation

$$\hat{Y} = 20.000 + .455(X) \tag{11}$$

as “the direct effect of the constant on endogenous variable *Y*” (p. 371). The “constant” is a 1.0 value placed behind the intercept, which permits rewriting the equation as

$$\hat{Y} = 20.000(1.0) + .455(X) \quad (12)$$

and on mid-page 370 as

$$\bar{Y} = 20.000(1.0) + .455(\bar{X}) \quad (13)$$

Unfortunately, it is incorrect to interpret the intercept 20.000 as “*the direct effect of the constant on endogenous variable Y*” (p. 371). One way to see why begins with noticing that the intercept depends on the scales of  $X$  and  $Y$ . If the scale for Kline’s  $X$  variable (Table 15.1, p. 370) had resulted in each case’s  $X$  score being 10 units higher, the  $X$  mean would increase by 10, and  $X$ ’s mean contribution in Equation 13 would increase by 4.55 units; the intercept would correspondingly decline by 4.55 units, from 20.000 to 15.45. This imagined change in  $X$ ’s scale alters the intercept *without* changing  $Y$ , the 1.0 constant, or any variable’s effectiveness—namely without changing *any* feature in Kline’s interpretation of the intercept as “*the direct effect of the constant on endogenous variable Y*”! Similarly, had  $Y$ ’s values been reported on a scale reading 20 units lower, Kline’s equation would become  $\hat{Y} = 0.0(1.0) + .455(X)$  and the “constant” would now seem to have no “effect” (according to Kline), even though this change in  $Y$ ’s scale does not change the causal effectiveness of anything.

But deeper causal issues are also involved. Consider the word “constant” in describing an intercept as “the direct effect of the *constant* on endogenous variable  $Y$ ”. Can a *constant* have effects, direct or otherwise? The answer is no—constants do not have “effects,” because effects demand potentially different outcomes resulting from *variations* in the cause (see Pearl’s definition; 2000: 70), or Mulaik’s insistence on variables (2009: 84–102)). Kline understands “the constant...is not a variable in the usual sense, because it has no variance” (p. 371), but he persists in referring to the intercept as the “effect” of the constant even though there is no such thing as an effect without potential variability in the cause. Placing a 1.0 after the intercept mimics the positioning of a causal variable like  $X$  but is insufficient to warrant interpreting the 20.000 intercept as an effect.

Next, consider that just as error variance changes upon addition of new predictors, intercepts also change upon addition of new predictors. And just as we do not know which real variables contribute error variations in equations, we do not know which real variables contribute to the intercepts in equations. Each equation’s error variable’s variance stands-in for the variation-consequences of unspecified causal variables, and each equation’s intercept stands-in to coordinate the effects and scales of all the unspecified causal variables, with the effects, scales, and means of all the included variables. Failing to acknowledge intercepts’ connections to real excluded variables leads Kline to the problematic claim that “If a variable is excluded from the mean structure, its mean is assumed to be zero” (p. 372). That is false because including a new clause adds a term containing the effectiveness and mean of that variable to the right of equations like Equation 13. A *regression* error variable is assumed to have a zero mean, but a real excluded causal variable need not have zero mean. If the  $Y$  variable in Equation 13 was exogenous because it had no specified causes, the  $Y$  mean would correspond exactly to the intercept but that would not force the unmodeled causes of  $Y$  to all have zero means.

Now consider that each different equation’s intercept, and each different exogenous variable’s mean, depends on a unique set of excluded variables having different scales, means, and degrees of effectiveness. Contrast this with **the constant** in Kline’s intercept interpretation, represented by **the triangle-containing 1** in Kline’s Figure 15.1 (as duplicated at the top of Figure 3). Kline’s figure represents **the constant** as the (singular) common cause of two variables, and as the common cause of many more variables in figures on pages 385, 389, 404, and 412. If the constant really was a common cause influencing two downstream variables, this would result in covariance between the two effected variables (recall Figure 1C and Equation 10 above). Unfortunately, Kline’s representa-



tion is inaccurate: the arrows connected to the triangle-1 are not effects, the triangle-1 is not a variable, and the implication of covariance would be wrong. Each endogenous and exogenous variable should be granted its own “constant” corresponding to the unique set of unmodelled variables contributing to the constant following/attached-to the intercept for each endogenous variable, or to the mean of each exogenous variable, as depicted lower in Figure 3. There the dotted lines are obviously not effects, because they have no arrowheads, and each variable is granted its own constant—whether representing an intercept or exogenous variable’s mean. This representation is similar to that employed by Hancock, Kuo, and Lawrence (2001), but the absence of arrowheads ensures these are not interpreted as “effects,” and deletion of the triangle-1 minimizes space requirements. Placing the intercept/mean designation near the variable’s unmodelled sources of variance (disturbance/error variance for endogenous variables; total variance for exogenous variables) signals that the unmodelled causal variables contributing variance also contribute to the corresponding variable’s mean or intercept.

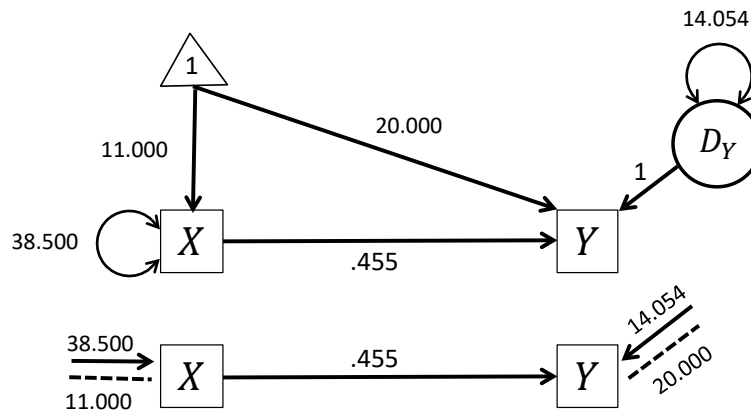


Figure 3. Kline’s Figure 15.1 (top) and a replacement (bottom).

The literature is not yet committed to a single felicitous mode of diagramming means and intercepts, but we can, and should, be consistent in our verbal descriptions. Consider part 2 of Kline’s Rule 15.3, which claims that “for endogenous variables, the direct effect of **the** constant is an intercept but the **total effect** is a mean” (p. 372; emphasis added). The intent of this “rule” is reasonable, but its execution is problematic. The intention behind “total effect” can be seen by writing the equation for  $Y$ ’s mean in Kline’s Figure 15.1 as

$$\bar{Y} = 20.000 + .455(\bar{X}) \tag{14}$$

and rewriting this with the numerical  $X$  and  $Y$  means, and inserting the constant (1.0):

$$25.000 = 20.000(1.0) + .455(11.000)(1.0) \tag{15}$$

This equation corresponds to the math on pages 370 and 372, and makes it *look like* the  $Y$  mean (25.000) is the sum of a direct “effect” of the (1.0) working through the intercept of 20.000 and an “indirect effect” of the (1.0), calculated as the produce  $.455(11.000)$ —so Kline expresses the  $Y$  mean as a total effect of the constant. The problem is that the intercept (20.000) and  $X$  mean (11.000) are not effects, and there really is not *one* single/lone constant. Each (1.0) refers to a *different* set of variables. In Equation 15, the left-most (1.0) refers to the disturbance/error/unavailable causes of  $Y$ , while the right-most (1.0) refers to entirely unknown causes of the  $X$  mean. The (1.0)s look the same, and the triangle-enclosed 1 in Kline’s Figure 15.1 looks like a single variable—but it isn’t. There are in fact multiple (1.0)s, and each refers to a different set of variables.

This recommends modifying Kline's statement quoted above to say that an endogenous variable's mean can be calculated as the "total effect" of *multiple different sets of variables*—which is glaringly self-contradictory, because total effects tabulate effects originating from a single source, not effects originating from multiple different sources. Put simply, the verbal reference to "*the constant*" inappropriately conflates multiple different causal entities, and recommends multiple corrections to Chapter 15 (see p. 371, 372, 378, 379, 380, 383, 384, 386, 387, 388) and Chapter 16 (p. 403, 420). The calculations of product terms work largely as Kline reports, but Kline's linguistic and diagrammatic descriptions of why things work that way require substantial revision.

In addition to the above, Kline's presentation of means and intercepts would benefit from observing that many social science variables have arbitrary scale-origins and scale-units. This would provide an opportunity to address the difficulties involved in locating non-arbitrary scale origins and units of measurement, as well as clarify why even overidentified SE models with mean structures have a limited ability to assist in locating non-arbitrary means and intercepts.

### **Kline's detailed example of means and intercepts**

Kline illustrates the modelling of means and intercepts using data from military personnel repeatedly attempting an air traffic controller exercise (p. 375–87). Kline bases his example on 137 cases that Browne and Du Toit (1991) selected for reanalysis from experiments conducted by Kanfer and Ackerman (1989). I suspect, but was unable to confirm, that the cases came from Kanfer and Ackerman's third experiment, and I was unable to determine how these cases had been selected from Kanfer and Ackerman's many cases, or even whether these cases came from an experimental or control condition. We cannot hold Kline responsible for Browne and Du Toit's failure to report how they selected their cases, but we should hold Kline responsible for emphasizing an example that precludes careful consideration of the data-gathering details that constitute the bedrock of competent structural equation modelling. As we shall see, this provides an instance of learning the hard way.

First, a caution. The relevant data matrix appears in Table 15.3 (p. 376) and is reported as based on  $N=250$  when in fact these statistics were based on  $N=137$ . Kline says some "technical problems" (p. 375) were avoided or resolved by *artificially* increasing  $N$  from 137 to 250, but he provides no indication of the nature of the resolved problems. Artificial increases in  $N$  are disconcerting and should be discouraged, but this seems likely a mere indiscretion in comparison to another feature of Kline's Figures 15.3 and 15.5 models that is likely to be unjustifiably emulated. Kline includes, but never defends, why these models permit correlation between successive (time-adjacent) measurement error variables. It is easy—too easy, and too easily emulated—to contend that the mere proximity of one measurement to the next warrants measurement error covariances. Omitting these error covariances results in model failure (Table 15.4), but even with the error covariances included, the models remain significantly inconsistent with the data. Kline "retained" both models, thereby persisting in his troubling disregard for model test evidence (p. 380, 385), but his inclusion of dubious error covariances to transform a highly data-inconsistent model into a model displaying modest but still significant ill fit enticed me to consider this more carefully.

I used LISREL to replicate the results reported in Kline's text and website, and then altered the model in a way I considered to be more theory-defensible. I thought each subject's real performance on the air traffic control learning task at any one time would influence their subsequent performance. That is, I viewed each participant's true and improving performance at any one time as likely to persist and contribute improved performance on their next attempt at

the task. This conceptualization recommends replacing Kline's five error covariances with five effects, leading from the performance on each trial to the performance on the next trial. The result was that my version of Kline's Figure 15.3 model fit ( $\chi^2=6.422$ ,  $df=7$ ,  $p=.492$ ), while Kline's model did not ( $\chi^2=16.991$ ,  $df=7$ ,  $p=.017$ ; p. 381); and my version of Kline's Figure 15.5 model fit ( $\chi^2=7.886$ ,  $df=11$ ,  $p=.724$ ), while Kline's model did not ( $\chi^2=27.333$ ,  $df=11$ ,  $p=.004$ ; p. 381). These  $\chi^2$  values use Kline's artificial  $N$  of 250, and my models' fits were further "improved" by using the proper  $N=137$ . I encountered no unusual estimation difficulties using the real  $N$ , and I encountered no sign of technical problems that might have warranted using Kline's artificial  $N=250$ .

My models closely reproduced the pattern of means in the data, and confirmed the relevance of the "Ability" variable in Figure 15.5, but provided somewhat different explanations for how the observed means are accounted for via Kline's Initial and Shape latents. Kline knows that in the context of latent growth curves, earlier observations of values of a variable can sometimes influence later values (see his p. 391 figure), though my models recommend retracting his claim that effects leading to successive observations and questions regarding latent growth curves "cannot be answered in the same model" (p. 392)—because that is precisely what my models did. I should also report that including measurement error variance in the Ability variable further improves my model, and would presumably improve Kline's Figure 15.5 model if he included measurement error variance on this single indicator.

The gist of this story is that Kline was caught in the act of inserting undefended error covariances merely to reduce  $\chi^2$  ill fit, when in fact a substantively reasonable and cleanly fitting model was easily attainable. The warning is clear: Do not insert coefficients merely to improve fit. The alternative is equally clear: Pay attention to the data-providers' causal world. Readers are also encouraged to consider why Kline failed to detect his problematic model specification despite employing the two-step approach that was supposed to make "it easier to detect potential specification error" (p. 376–77).

## **Chapter 16, Multiple-samples analysis and measurement invariance**

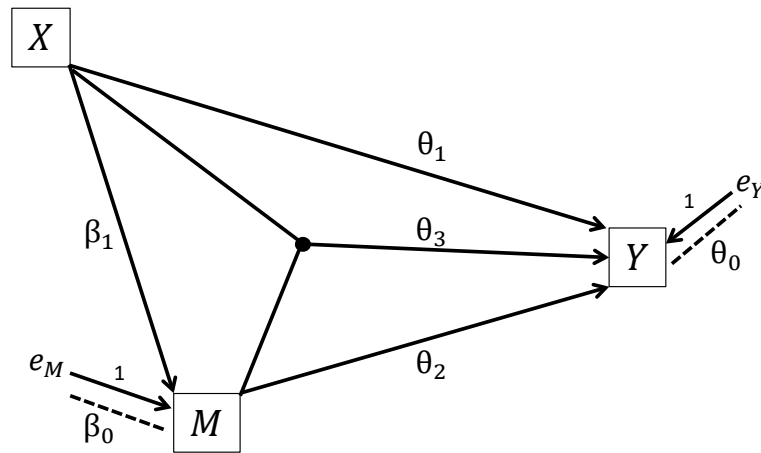
Multi-group analyses are most commonly used to assess measurement invariance, which Kline illustrates using both continuous and ordinal indicators. Additional uses for multi-group analyses are granted a paragraph of discussion (p. 395) but readers are not informed about how such models can be used to: identify otherwise underidentified models, control for variables that are unmeasured in some groups, or to integrate complementary but non-overlapping model segments from diverse data sets (Hayduk 1996: Chapter 5).

As in the preceding portions of Kline's text, the writing is effective and informative but also laced with model-testing laxity and the occasional oddity. Kline's section on "testing strategy and related issues" reports that "Failure to retain the invariance hypothesis at a particular step means that even more restrictive models are not considered" (p. 399–400), but Kline fails to explicitly report that this testing requirement begins at the very first step (namely, with the baseline configural model) because  $\chi^2$ -difference testing (such as for equality of coefficients between-groups) is only statistically justified if the sampling distribution for the less-constrained model has a  $\chi^2$  distribution. Hayduk (2016) underscores this point and illustrates a way to strengthen baseline configural model assessments. Fortunately, Kline's exemplified baseline configural model fits the data. An oddity is that Kline rejects this fitting model and adds a coefficient (on the basis of an ill fit covariance and without substantive justification) to further improve the model fit (p. 406, Table 16.2 and website). Kline's next example also inserts a coefficient merely to improve fit but reverts to "retaining" failing models (p. 416–17, Table 16.6).

## Chapter 17, Interaction and multi-level modelling

This chapter begins with an effective presentation of interaction with observed variables, but continues Kline's waffling between regression models and causal models (p. 424–29). Causal requirements emerge by page 432, but Kline's unwillingness to routinely encourage and require causal structuring becomes uncomfortably obvious when later on that page he applies the term "cause" when time-sequencing is available and "effect" for cross-sectional designs. Unfortunately time sequencing is not an appropriate criterion because reciprocal effects can be estimated with cross-sectional data and both the reciprocally connected variables can't possibly be "first." Urging use of the terms *cause without effect*, and *effect without cause*, along with a dubious differentiating criterion, could be called self-inflicted befuddlement.

My following comments aim to improve the interpretation of mediated interactions (as discussed on p. 435–37) by expanding Kline's Figure 17.5c example to illustrate some general principles and important overlooked complexities. Figure 4 corresponds to Kline's Figure 17.5c, with coefficients from his Equation 17.7. Intercepts are represented with dashed lines and disturbances/errors are designated as  $e$ 's. Kline expressed the model as "two unstandardized regression equations" (Equations 17.7; p. 435, emphasis added). I express these as putatively causal equations by replacing the regression-predicted  $\hat{Y}$  values with  $Y$  and the relevant causal-disturbance/error  $e_y$ , and by similarly converting the mediator-moderator "regression" equation into a causal equation by replacing  $\hat{M}$  with  $M$  and disturbance/error  $e_M$ .



**Figure 4.** Kline's Figure 17.5c, expanded and re-expressed with coefficients.

$$M = \beta_0 + \beta_1 X + e_M \quad (16)$$

$$Y = \theta_0 + \theta_1 X + \theta_2 M + \theta_3 XM + e_Y \quad (17)$$

The first model equation reports  $X$  as a cause of mediator/moderator  $M$ . Since  $M$  causes  $Y$  in the second equation,  $M$  is postulated as mediating part of  $X$ 's causal impact on  $Y$ .  $M$  also functions as a moderator (interacting variable), represented by  $M$ 's product with  $X$  in the second equation.  $\beta_0$  and  $\theta_0$  are intercepts capturing the net impact of variables currently excluded from the equations. Before returning to Kline, we rearrange these equations by inserting the first equation into the second (which corresponds to replacing  $M$  with  $M$ 's causal foundations):

$$Y = \theta_0 + \theta_1 X + \theta_2 (\beta_0 + \beta_1 X + e_M) + \theta_3 X (\beta_0 + \beta_1 X + e_M) + e_Y \quad (18)$$

and multiplying out while inserting { } to keep track of the origin of the various terms:

$$Y = \theta_0 + \theta_1 X + \{\theta_2 \beta_0 + \theta_2 \beta_1 X + \theta_2 e_M\} + \{\theta_3 X \beta_0 + \theta_3 X \beta_1 X + \theta_3 X e_M\} + e_Y \quad (19)$$

Each term in this equation provides a causal constituent of  $Y$ , and the bolded terms document where and how the  $X$  variable participates in producing  $Y$ .

[Supplement Section 6](#) considers each right-hand term to determine what the model claims would be the consequences of an intervention changing a treatment  $X$  from 0 (no treatment) to 1 (treatment). Each term in the expanded equation is examined to see whether and how it responds to the postulated intervention. The relevant interpretation consists of whatever wordings accurately describe the right-hand terms changing as a consequence of  $X$  changing from 0 to 1, and the composition of each term details the causal features providing that component of the effect transmitted to  $Y$ . The required assumptions consist of wordings reporting features required to render some right-hand terms constant and hence unable to produce change in  $Y$  as  $X$  changes from 0 to 1. A model's implications for a postulated intervention have been thoroughly considered if the researcher examines the coefficients and variables comprising all the terms in  $Y$ 's expanded equation, and reports the assumptions/presumptions required to attain and respect the intervention of interest.

Kline does not report equations corresponding to Equations 18 or 19 and instead moves directly from the model equations (like Equations 16, 17) to a set of equations reporting how to calculate the effects of intervening to change  $X$  from 0 to 1. [Supplement Section 6](#) follows the procedure outlined above and locates some unacknowledged assumptions and requirements of Kline's effect calculations. For example, Kline's equations are inappropriate if the treatment happened to be coded 1 = no treatment and 2 = treatment instead of 0 and 1, and his equations do not apply if there are two treatment levels so that 0 corresponds to no treatment, 1 to weak treatment, and 2 to strong treatment. And Kline's formulas apply only to this specific model and one specific intervention. The [Supplement Section 6](#) procedure of beginning with the equation for the dependent variable of interest, and replacing each moderator variable in the equation with that moderator's causal sources, is applicable to a wide variety of models and can examine a diverse range of potential interventions.

As it stands, researchers with more complex conditional models are cornered into trying to squash descriptions of their model into Kline's wordings rather than having been equipped to develop interpretations appropriate for their particular model. Kline's text leaves readers ill-prepared for considering consequences of interventions unavoidably making two simultaneous changes, or the consequences of reducing or increasing some effect (without intervening to change any variable), or assessing which specific disturbance/error variable's values might disrupt or assist the causal impact of interest. Assessing such interventions becomes feasible using the procedure illustrated in the [Supplement](#).

Just as the interaction/moderator segment of Chapter 17 could be improved by considering model equations (as above), the chapter's discussion on multi-level modeling could be similarly improved. Kline's figures, Mplus syntax, and output (on the publisher's website) are appropriate but the basic principles and model details remain obscure without the model equations. Equations would clarify why  $s_1$  and  $s_2$  appear in one portion of Figure 17.7c as effects and in another portion as variables. And model equations would clarify why "Game" is boxed in one portion of the figure and circled in another. Currently the reader is left puzzling how slopes and intercepts can be variables, and can be justifiably perplexed by noticing slopes are designated  $s_1$  and  $s_2$  while the intercepts seem to be AWOL. How is a reader to understand why the same indicator variables appear in two parts of Figure 17.8b, and determine whether the disturbances/errors on the duplicate indicators are the same? Clearly there are too many potholes for this review to fill, though the road to improvement is paved with equations.

## Chapter 18

Kline's concluding chapter accumulates and effectively structures the recommendations and advice provided in earlier chapters, and hence it reflects *Practices* but not "*Best Practices*" in *Structural Equation Modeling*. The chapter begins by tabling several references offering suggestions on conducting and reporting SEM studies. The table's footnote reports the third of ten SEM commandments as: "simpler models are better" (p. 453). A simple model of a moderately complex world is likely to be misspecified, so surely "Best Practice" would recommend an appropriately complex model, not merely a simpler model!

"Best" requires acknowledging that some practices are better than others, and support for the stronger practice. Unfortunately, if we consider model testing, Kline continues to promote weak practice. Model testing is not even granted its own section in Chapter 18, and it is mentioned as only one of 16 points under the heading *Estimation*. Even there the wording "Never retain a model based solely on global fit testing" (p. 461) is slanted to suggest retention of test-failing models, as opposed to respecting evidence and pursuing the sources of detected model-data inconsistencies.

Kline's Chapter 18 sections on model *Specification* (p. 454) and *Respecification* (p. 463) could similarly be strengthened by encouraging consistent pursuit of models mirroring the world's causal structure. Structuring models to reflect specific theories is laudable but limited. A researcher committed to a theory-based model that demonstrates data-inconsistency will flounder until they re-ground themselves in the quest for understanding the world by seeking a new or modified theory. Consider the risk created by routinely including residual/error correlations in models (recall the problematic models from Chapter 15, p. 378, 385), and the risk arising from attempting to fix failing models by adding coefficients suggested by modification indices or specific residuals (recall the problematic negative estimate p. 347). The risk is *not* merely of "capitalizing on chance" (p. 455), or that this constitutes an exercise in "chasing sampling error" (p. 463), or that this incurs a "cost of too many parameters" (p. 463). This risks obscuring (by incorrectly modeling) real stable evidence that is inconsistent with the substantive structure of the current model. Inserting coefficients merely as a matter of "routine" or because the coefficients have large modification indices risks inserting worldly-inconsistent coefficients that absorb and obscure whatever real data covariance inconsistencies managed to speak against the original theory/model. The fundamental risk is that real (not merely sampling error) covariances are modeled in the wrong way. Replication will not detect or correct models based on improper "causal" accounts of real covariances. The real covariances remain stable and so the researcher is condemned to proceeding with a now-fitting-and-replicating but nonetheless wrong model, and hence is robbed of the opportunity to get the model right.

Lack of commitment to seeking the world's structure is also evident when Kline says "do not specify feedback loops as a way to mask uncertainty about directionality" (p. 455). The options Kline leaves open to this researcher are to choose one causal direction or the other, or to drop both effects. That is, Kline implies the researcher should include a non-theory based effect-directionality (or gap) into their model, rather than encouraging the researcher to introduce exogenous causes that would make the reciprocal effects estimable and thereby permit the worldly-data to potentially support the existence of both, either, or neither of the theory-eluding reciprocal effects.

Similarly, consider the flaccid commitment to seeking a world-matching model in the context of measurement. "Multiple-indicator measurement is generally better than single-indicator measurement" (p. 454). If the researcher begins by providing each latent the best available indicator, each additional indicator is prone to being weaker and more problematic, and hence more indicators does not necessarily constitute better modeling. "Best practice" would begin with the best indicators and supplement these with only strong additional indicators. Two or three indicators

per latent substantially increase model testing power (Hayduk and Littvay 2012) and are likely to be sufficient to detect specification problems—presuming the researcher respects model-test evidence of problems. Multiple indicators unavoidably include weaker indicators, and expand models in ways which tend to squeeze out latents clarifying mechanisms of action or contributing informative controls—which results in generally worse, not better, models.

And consider the claim that a way to improve on a single indicator is “to specify an instrument for the single indicator” (p. 454). A reader would be justifiably mystified by how a problem with a single-indicator is to be overcome by introducing another single-indicator (namely the instrument) into the model. By downplaying the relevance of the world’s causal structuring, Kline is cornered into expressing this as if the improvement somehow comes from statistics (the statistics of instrumental variables) rather than from employing single indicators in ways that benefit from, and capitalize on, the world’s causal structure.

Turning to *Identification* (p. 457), it is reasonable to check that the number of data covariances exceeds the number of estimated coefficients, and the identification of simple models should indeed be checked. But the unavailability of general procedures for checking full or moderately complex models should have prompted suggestions for: locating likely problematic model segments, checking maximum likelihood iterations (if maximum likelihood estimation is used), checking for unexpected estimate signs or magnitudes, and checking for inflated standard errors. Solutions to underidentification should also have been included—namely adding data constraints (e.g. additional identification-helpful variables) or adding model constraints (e.g. fixing/specifying or constraining model coefficients). Archival data may offer fewer opportunities to improve identification by adding relevant variables, but it is simply a mistake to claim “that the model is not identified” (p. 459) merely because it is based on archival data. Fixing/specifying coefficients to attain identification is particularly relevant with archival data, especially if the researcher investigates the sensitivity of the model to a realistic range of fixed coefficient values—including non-zero values for unresolved latent-level loop or reciprocal effects.

Model *Respecification* (p. 463) provides another instance where it would be helpful to differentiate between fitting models and proper models. Consider a researcher in a discipline confronting worldly causal structures that are not yet understood. If the researcher’s model fails, the basic options are: add coefficients that reduce the model’s ill fit, report model failure, or junk the model. Junking the model seems a waste, and reporting failure of a model is likely to be personally uncomfortable, so the common response is to add coefficients until the model’s fit can be passed off as good enough. Unfortunately, as long as model respecification is touted as being a matter of each particular model’s local or global fit, the respecification is likely to fall short of addressing the deeper disciplinary concerns. The concern is not that the “good fit is achieved at the cost of too many parameters” (p. 463). The concern is that even one additional coefficient may be sufficient to obscure the evidence recommending that the discipline reconsider the thought modes underlying the whole model.

## Your first edition, or Kline’s fifth edition

I have been unable to convince myself of the source(s) of Kline’s reticence to notice and address the multiple and diverse concerns discussed above. To see how Kline thinks about these matters, and to glean a hint of his intentions, I requested that the editor of *Canadian Studies in Population* (Frank Trovato) invite Rex Kline to respond to this review essay. It would be nice if Kline plans a fifth edition, but if this is not planned, I hope that readers will consider preparing their own first edition, or possibly one co-authored with Kline. Irrespective of who writes the next edition, I would suggest a title like *Principles Nurturing Best Practice in Structural Equation Modelling*, where the

text begins by focusing on structural equation models as striving for correct causal representations (a commitment which differentiates SEM from regression) and complementing this with routine attention to detecting and resolving model misspecification (not merely seeking fitting models). Whether or not you are the new-author, you can do SEM a service by inserting a reference to this review in whatever copies of Kline's fourth edition you encounter.

## Acknowledgement

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## Supplement to review essay on Rex B. Kline’s *Principles and Practice of Structural Equation Modeling*

Leslie Hayduk

This supplement provides contextualizing discussions for several sections of the review essay, and ends by listing editorial corrections to Kline’s text.<sup>1</sup>

### Section 1: Causal structures versus regression equations

*Finding causal action:* Does real “isolation” (p. 27) require real causal isolation? Is “independence of residuals and predictors” (p. 27) founded in real causal separateness? Kline says regression equations make “less sense” if they misrepresent the causal world (p. 27) but why only “less sense” rather than SEM-nonsense? Are regression predictions likely to hold if some regression-predictors happen to be effects rather than causes? What is the nature of explanation and explained variance (p. 29) if the supposed-explanatory variables are not causes? What if not causal actions account for why regression estimates change with introduction of control variables (p. 32)? Kline describes the “statistical and conceptual assumptions of regression” (p. 33) as including that “*there are no causal effects among the predictors (i.e., there is a single equation)*” (p. 34; emphasis in the original). This is false. Regression does not *assume* there are no causal effects among the predictors, even though it does not estimate potential effects among the predictors in one equation. (A regression equation paralleling Equation 5 would acknowledge and adjust for the covariance between the two causes, but would leave unspecified whether the covariance arose from either of the causes influencing the other.) What produces bias in regression estimates (p. 35) if not misrepresentation of worldly effects? Isn’t a “serious specification error” (p. 35) a causal error? Kline downplays the causal basis of several matters by repeatedly presenting how the numbers and equations work out, rather than attempting to explicate how *proper* representations and *mis*-representations of the world’s causal structuring lead to the statistics working out as they do. The overlap in Kline’s Venn diagram (p. 40) could be usefully connected to the covariance term in essay Equation 7 above, but his couching this in a discussion of part- and partial correlations and regression  $R^2$  values (p. 39–41) disconnects it from structural equation models. (A similar comment applies to Equations 2.13 and 2.14.)

Kline’s timid differentiation between SEM’s causal-focus and regression’s causal-indifference is sprinkled throughout the text and appears in many guises. For instance, Kline’s example of left-out error variables (p. 36) would be easier to understand if the substantial correlations were characterized as originating in overlooked causal connections, and if “relevant predictors” (p. 36) had their relevance grounded in causal actions, rather than leaving the foundations of relevance unspecified—or, worse yet, permitting readers to incorrectly presume that higher correlation constitutes or justifies the relevance of a “predictor.” For another example, “Suppression” (p. 36–37) naturally and easily meshes with the causal understanding that some effects may counteract other effects. Isn’t “suppression” just another name for counteracting causal actions? Kline could not entirely avoid appealing to “indirect effects” as a foundation of suppression—though he does manage to delay it until the section’s second-last sentence (p. 37).

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1. Cited references are listed at the end of the Review essay.

Additional connections appear in the discussion of indicators in factor analysis, where Kline says “Common variance is shared among the indicators and is a basis for observed covariances among them” (p. 190). Actually, a latent factor functions like the common cause in Figure 1c, and gives each indicator its own variance (via essay Equation 3), and produces covariance among each pair of indicators though causal actions (via essay Equation 10). There is no sharing of variance—each indicator has its own. And consider Chapter 13 on Confirmatory Factor Analysis Models (CFA models). According to Mulaik, in doing CFA “the researcher begins with a conception of a set of latent exogenous **causal** variables having specified **effects** on a set of endogenous manifest ‘indicator variables’” (2009: 219; emphasis added). In factor analysis, the effects of latents on their indicators are historically called loadings or pattern coefficients, and Kline says, “Pattern coefficients are interpreted as regression coefficients” (p. 301)—where regression’s causal indifference is clearly inconsistent with Mulaik’s emphasis on cause! And consider “reification”: it would indeed be an error to claim an underlying latent factor “*must* correspond” (p. 300; emphasis in the original) to some real thing because a model can be wrong; but it is not an error, and actually is SEM-positive, to strive to make SEM latent variables correspond to real worldly entities. It is not just that models are “most meaningful” (p. 306) when they are consistent with the data provided by the causal world—they lose meaning if the model’s structure fails to correspond to the world’s structure. And causal actions explain why different model coefficients can have nearly identical modification indices (p. 312)—this happens when two currently omitted coefficients have nearly identical causal implications for the covariances between the observed variables. And notice that Chapter 10 considers “Structural Regression Models,” as if the latent-level structures are regression, not causal, structures. See page 232, where the effect in a path model striving for causal correctness is supposed to be “interpreted exactly as a regression coefficient,” despite regression not striving for causal properness.

Similarly, in Chapter 14 we read: “path coefficients are interpreted for SR models as **regression coefficients** between factors. Total **effects** between factors can be decomposed into direct and total indirect **effects**, just as in path analysis” (p. 340; emphasis added). Are regression coefficients really effects? Kline stumbles repeatedly over whether or not the latent levels of structural equation models strive for causal standing.

Chapter 17 on interaction and multilevel modeling begins with regression and yet causal effects soon appear in the text and section titles (p. 427, 431). By page 432 we are told that “just as a mediational model is a causal model, so too is a model of moderation”—but the location of the transition from regression to causal action was left a mystery. Kline’s artificial data illustrating this chapter’s introductory “regressions” (p. 424–29) actually has a known causal foundation—one that is not reported, and is sloughed over. Then, consider moderated mediation, where the “interactive effect is represented in the figure by the *regression* of  $M$  on  $X$ ,  $W$ , and  $XW$ ” (p. 434; emphasis added). Does the estimate really constitute an effect if the equation is not causal and merely a regression? The delay in introducing required causal action robs Kline of yet another opportunity to instruct his readers on the necessity of attending to causes *before* specifying interactions or multilevel models, and robs him of the opportunity to instruct readers to check causal specifications both before and after running models that contain interactions or multilevel effects.

## Section 2: Reciprocal effects

Here is a difficulty produced by Kline’s omission of reciprocal effects. We are told that “within-time associations in panel models are typically specified as unanalyzed” (p. 139), and Figure 6.8b complies with the absence of within-time causal arrows. Unfortunately, this results in a seemingly unnoticed logical difficulty. Consider just the  $X$  row and  $M$  row of Figure 6.8b (p. 140), and see

the absence of an effect of  $X_{12}$  on  $M_{12}$  and the presence of an indirect effect  $ec'$  connecting  $X_{11}$  to  $M_{13}$ . The model precludes  $X$  from influencing  $M$  at any specific time, yet permits  $X$  to effect  $M$  by skipping from some earlier time to some later time. It is logically awkward to claim that  $X$  does not cause  $M$  at any specific time while simultaneously claiming  $X$ 's causal actions somehow jump to causing  $M$  at a later time. The past is influencing the future without ever going through the present? How small (but not infinitely small) time frames for “the-present” connect “the-past” to “the-future” is a matter well worth considering, especially since SEMs need *not* be confined to modelling causal actions occurring instantaneously at the indicator-observation times (Hayduk 1985, 1996).

### Section 3: Separating observed from latent variable

Here are some “unusual” features of Chapters 6 and 7, originating in their focus on observed-variable path models, despite almost all measured variables containing some measurement error. The rarity of error-free variables probably explains why the only real example in Chapters 6 and 7 is a seven-line discussion on page 159, and even this example ignores likely measurement errors in the relevant modelled variables. The points in these chapters were obviously made with fictional examples, and could have been made as easily, and more appropriately, using fictional latent-variable examples. The artificiality of purely “Observed-Variable (Path) Models” will be obvious to anyone noticing that Chapter 7 begins by reporting “two general...requirements” for such models, the second requirement being that “Every *latent* variable...must be assigned a scale” (p. 145; emphasis added). Inserting required features for latent variables in observed-variable models is probably a consequence of Kline's chapter reorganization, but at least this moves latents toward where they should have been all along.

### Section 4: Some nit-picking concerns connected to Chapter 8

To my knowledge, there has been no demonstration of an appropriate way to address the multiple testings of independencies (and manufacturing of multiple dependencies by controlling for colliders) derived from DAG investigations (e.g., p. 172). Another feature for which I know of no current resolution concerns the “basis set” of non-redundant conditional independencies (p. 173). Independencies beyond the minimal set are redundant if the independencies hold, but if one or more do not hold, that raises the possibility of dredging diagnostic assistance from the initially declared-redundant set. DAGs also await ways of appropriately assessing fixed model coefficient values, and differentiating between properly and improperly fixed/specified coefficients. Kline cannot be held responsible for these loose ends in the DAG literature, but a caution should at least report that various extensions remain to be pursued. What Kline can be faulted for is his failure to integrate the new DAG testing precision with his discussions of testing in Chapters 11 and 12.

### Section 5: A passel of technical teasers connected to deficient testing

Regarding page 60, paragraph 2, second line: is the null hypothesis of the *model* test really a hypothesis that is “false by the degree indicated” by some index? I vote no, it is not.

Regarding page 239, point 2: the error rate might be high (for some unspecified test, for some unspecified conditions, and without reference), but what kind of error rate would be typical for reasonable conditions?

Regarding page 239, fourth-last line: if the global test indicates model misspecification, the coefficient estimates are likely to be biased. Would you want to pay more attention to the biased estimates?

What constitutes the “population” in model testing? In  $\chi^2$  testing the  $\Sigma$  matrix, namely the model-and-estimate implied covariance matrix is employed as the population. This may or may not correspond to the “worldly population” that provided the data, and it is not dependent upon the data coming from a random sample of any “worldly population.” Several of Kline’s wordings about these distinct population are sufficiently ambiguous as to invite misinterpretation (p. 235, 236, 265), as well as confusion about the nature of the “multivariate normality” and the hypothetical “random sample” (p. 270) relevant for maximum likelihood estimation and  $\chi^2$ .

The  $\chi^2$ -difference test is justified only if the “more complex of the two models compared” (p. 281) actually fits the data, and not merely that it should “fit the data reasonably well” (p. 281, and see p. 306).

“Given two models with similar fit to the data, the simpler model is preferred, assuming that the simpler model is theoretically plausible” (p. 128)—so the model is “preferred” even if both models are similarly highly inconsistent with the data!

A perfectly fitting model may not be perfect because seriously misspecified models can fit perfectly (Hayduk 2014a). And there is no known way to calibrate the amount of ill fit to the seriousness of the corresponding model causal misspecification—not even with a noncentrality parameter. So what is the nature of the “perfection,” and what makes some amount of ill-fit “acceptable” when Kline says, “what is considered ‘acceptable’ departure from perfection is related to the value of the noncentrality parameter for the  $\chi^2$ ” (p. 60)? The size of a claimed-acceptable noncentrality is actually an index of the degree of shamefulness in the researcher’s disregard for evidence. Zero-noncentrality with zero-shamefulness is as easy as using the ordinary  $\chi^2$ .

Kline cites references that discuss ways SE model-testing differs from coefficient-testing (Hayduk 1996, 2014a; Hayduk and Glaser 2000), so it is surprising that his discussion of “Cognitive errors in significance testing” (p. 55–56) fails to distinguish between model tests and coefficient tests. I would have hoped that Kline would see that the testing differences also make it incorrect to treat confidence intervals around fit indices the same as confidence intervals around coefficient estimates. The difference between model adequacy and model fit means that if model “respecification is driven entirely by empirical criteria such as statistical significance, the researcher should worry—a lot, actually—about” *model misspecification* and not just “capitalization on chance” (p. 283). The issue that killed “automatic modification” was not capitalization on chance (p. 283); it was that ill-fit is prone to being inappropriately reduced by inclusion of additional *misspecified* coefficients matching *real* (not merely chance) residual covariances.

Overall, Kline could have avoided multiple imprecisions by acknowledging SEM’s commitment to seeking causal-theory, and emphasizing attention to evidence signaling the causal solidity of some models and causal dubiousness of others.

Rex Kline was one of the authors of a recent American Psychological Association publication which provided new reporting standards for structural equation models (see Table 7 in Appelbaum et al. 2018). I provided the following two replacements for sections of these standards to the SEMNET listserv (26 January 2018). Interested readers might see SEMNET for Kline’s response.

Replace the first sentence in the **Abstract** section with:

*Report a test of whether or not the model is consistent with the data, and the implications of informative localized-ill-fit.*

Change the title from Model fit to **Model Testing** and replace the first bullet point in this section with:

*Report the most powerful model test—usually chi-square (possibly adjusted) with its degrees of freedom and probability. All fit indices (even with values commonly reported as acceptable) are deficient at detecting model misspecification and hence cannot replace or displace the evidence provided by model testing.*

## Section 6: Improving Kline's Figure 17c example

The model-implied consequences of an intervention at  $X$  can be investigated by observing how the terms on the right of essay Equation 19 change in response to the postulated intervention. The model claims the effect of an  $X$  intervention on  $Y$  will be the difference between the  $Y$  value provided by this equation before the intervention and the  $Y$  value after the postulated intervention. Any right-hand terms left unchanged by, held constant by, or precluded by the imagined/postulated intervention will contribute equally to  $Y$ -before and  $Y$ -after intervention, and hence will not contribute to changing  $Y$ . The terms that are changed as a result of a postulated intervention document the components of the causal system contributing to changing  $Y$ 's value.

Kline postulates an intervention in which  $X=0$  constitutes a control condition and  $X=1$  denotes provision of a treatment (p. 435). (Readers familiar with Pearl (2000) will recognize the parallel to Pearl's *do(x)*.) Providing the treatment (namely, shifting  $X$ 's value from 0 to 1) in a world structured like essay Figure 4 would have two basic ways of influencing  $Y$ —namely, directly (with whatever complications moderation/interaction implies) and indirectly through  $M$  (also with whatever complications moderation/interaction implies). To minimize space, we will focus on understanding/interpreting the *indirect* effect on  $Y$  of an intervention changing  $X$  from 0 to 1. The focus on indirectness requires retaining terms containing both  $X$  and  $X$ 's  $\beta_1$  effect on  $M$ , but we must consider all the terms in Equation 19, which we duplicate here for convenience.

$$Y = \theta_0 + \theta_1 X + \{\theta_2 \beta_0 + \theta_2 \beta_1 X + \theta_2 e_M\} + \{\theta_3 X \beta_0 + \theta_3 X \beta_1 X + \theta_3 X e_M\} + e_Y \quad (19)$$

Now consider each right-hand term.

$\theta_0$  is the intercept corresponding to the net impact of variables not currently in the  $Y$  equation, which our “postulated intervention” did not address. This term forces clarification of our postulated intervention by demanding we add the assumption/pre-sumption that the excluded causes of  $Y$  remain constant and hence do not contribute to changing  $Y$  as the intervention changes  $X$  from 0 to 1.

$\theta_1 X$  is the direct effect of  $X$  on  $Y$  but we are seeking effects connected to, or functioning through variable/mechanism  $M$ , so we must clearly assert that our postulated intervention specifies this effect has somehow been rendered inoperative.

$\theta_2 \beta_0$  neither the  $\theta_2$  effect nor the  $\beta_0$  intercept are altered by the postulated change in  $X$  from 0 to 1, so this term remains constant, and hence does not contribute to a change in  $Y$  by the intervention of interest.

$\theta_2 \beta_1 X$  involves the indirect pathway due to the presence of both  $\beta_1$  and  $\theta_2$ , and this term's contribution to  $Y$  will change from  $\theta_2 \beta_1(0)$  to  $\theta_2 \beta_1(1)$  as the treatment changes  $X$  from 0 to 1, so this term contributes to changing  $Y$  by the intervention of interest.

$\theta_2 e_M$  our postulated  $X$  intervention and the causal action working through  $M$  did not change either  $\theta_2$  (the basic effectiveness of  $M$ ) or any unknown sources of the mediator-moderator  $M$  (namely,  $e_M$ ), so this term remains constant. This term instructs us that it would be possible to trace the consequences of an intervention changing the causal effectiveness of one variable at influencing another (like changing  $\theta_2$ ) rather than seeking the consequences of changing the value of a variable (like  $X$ ), but our particular intervention did not introduce such a change.

$\theta_3 X \beta_0$  This term will prove to be contentious, and I will return to it momentarily.

$\theta_3 X \beta_1 X$  involves an indirect effect of  $X$  through  $M$  due to  $\beta_1$ , and this term changes due to our postulated change of  $X$  from 0 to 1.

$\theta_3 X e_M$  This term is also contentious, and I will return to it momentarily.

$e_Y$  is rendered constant by again increasing the precision of our postulation, namely by adding the assumption/presumption that the intervention of interest does not alter any of the unknown causes of  $Y$ .

Now we return to Kline and notice the third Equation in 17.8 (p. 436) reports what Kline calls the Natural Indirect Effect (NIE) of an  $X=0$  to  $X=1$  intervention in this model as

$$NIE = (\theta_2 + \theta_3) \beta_1 \tag{S1}$$

Combining the two terms containing  $\beta_1$  highlighted above, and writing this as an effect (namely as an effect appearing in front of the  $X$  variable) results in something similar but not quite identical to Kline’s NIE equation.

$$[\text{indirect effect of}] X = [(\theta_2 + \theta_3 X) \beta_1] X \tag{S2}$$

$$[\text{indirect effect}] = (\theta_2 + \theta_3 X) \beta_1 \tag{S3}$$

The difference is the “extra”  $X$  inside the parentheses. If  $X$  moves from 0 to 1, both Kline’s Natural Indirect Effect coefficient (Equation S1) and our indirect effect coefficient (Equation S3) report exactly the same change in  $Y$ ’s value, because both provide zero contribution when  $X = 0$  and contribute  $(\theta_2 + \theta_3) \beta_1$  when  $X = 1$ , because the  $\theta_3$  is left unchanged by being multiplied by the “extra”  $X$  of 1 in Equations S2 and S3. A difference would arise if we had been considering a progressive  $X$  treatment, where a subject might receive partial treatment or multiple doses of the treatment, so  $X$  might take values like 0, 0.5, 1, or 1.5. If this is possible, our indirect effect calculation differs from Kline’s because the “extra”  $X$  in our formula alters the  $\theta_3$  portion of the effect (Equation S2 or S3). This cautions that Kline’s calculation of NIE applies only to dichotomous  $X$  variables scaled 0–1, not to  $X$  variables having non-dichotomous scalings, and not even to dichotomous interventions scored 1–2 rather than 0–1.

Now consider the terms skipped above. The value of the  $\theta_3 X e_M$  term, and hence  $Y$ , would change as  $X$  switches from 0 to 1 due to the postulated intervention if both  $\theta_3$  and  $e_M$  are non-zero. The disturbances or errors in equations are routinely presumed to average zero, but this equation contains each particular individual’s precise error/disturbance value not the average of multiple cases’ errors or disturbances (just as  $X$  and  $Y$  in the equation refer to specific, not averaged, values). This has two consequences. First Kline should have reported either that his NIE calculation (p. 436) presumes or assumes he is seeking the indirect effect for a case having precisely a zero disturbance/error, or that his calculation acknowledges the indirect effect will differ between cases and he is seeking only the average of those cases’ indirect effects. Second, Kline could have reported the possibility of assessing the extent to which variations in  $X$ ’s indirect effect on  $Y$  originates in variations in  $M$ ’s disturbance/error variable. The variance of  $M$ ’s disturbance/error is routinely estimated in structural equation models and hence the standard deviation of the disturbance/error is available. Calculating the magnitude of  $\theta_3 X e_M$  for error values one or two standard deviations above and below zero error would report the fluctuations in the postulated indirect effect likely to arise from the 0–1 change in  $X$  combining with modest or nearly-extreme disturbance/error values.

The consequences of the other skipped term, namely  $\theta_3 X \beta_0$ , are more awkward but intriguing. Here too, if  $X$  changes from 0 to 1, and if both  $\theta_3$  and  $\beta_0$  are non-zero, this term alters  $Y$ 's value in a way which involves  $M$  because  $\beta_0$  is part of  $M$ 's equation. Above it was relatively straight forward to presume  $e_M = 0$ , or equals zero upon averaging, to eliminate the term, but a corresponding presumption of  $\beta_0 = 0$  would be dubious. The  $\beta_0$  intercept is not an average. It possesses only a single value that is included/operative for each and every case, and like all intercepts its value depends on the scale and effects of all the variables included in the equation and even the scales and effects of excluded variables.

In what sense are the  $\theta_3 X \beta_0$  and  $\theta_3 X e_M$  terms “effects”? Neither term contains an “effect” if by effect we mean a coefficient depicting a regression-style slope, but both terms originate in  $M$  acting as a moderator variable that influences, changes, or adjusts the causal effect of variable  $X$  as it proceeds to impact  $Y$ . The term containing  $e_M$  describes effect-variations explicitly produced through the actions of unobserved causes that differ between individuals. The term containing  $\beta_0$  describes effect-variations produced through features that the model does not permit to vary between individuals, and that are partially controlled by both the zero-point and scale-units of the  $M$  variable (which are likely arbitrary).

And in what sense are the  $\theta_3 X \beta_0$  and  $\theta_3 X e_M$  terms “indirect” or “direct”? Tracing these terms back to Equation 19 finds that these originated in the multiplicative term connecting  $X$  and  $M$ , namely  $\theta_3 X M$ , and that the changes in these terms result from  $M$  responding to the postulated 0–1 change in  $X$ . Thus these terms clearly implicate the mediator/moderator variable, and can lay claim to being indirect consequences of an  $X$  intervention working through the mediator/moderator variable  $M$ . The  $\theta_3 X M$  term does not indicate whether  $M$  is modifying  $X$ 's effect, or if  $X$  is modifying  $M$ 's effect (namely whether the curvature in Kline's Figure 17.2 should be considered as changing slopes paralleling the  $X$  axis, or the other axis). Viewing this term as  $X$  conditioning/altering  $M$ 's effect makes the term seem like an indirect effect working through  $M$ . Viewing the term as  $M$  conditioning/altering  $X$ 's effect makes the term seem more like a direct effect. Kline, following Valeri and VanderWeele (2013), includes the term as part of what they call the “natural direct effect” NDE (p. 435–436, Equation 17.7) but that placement is debatable, and is better viewed as open to the researcher's preference for how this term would be most usefully considered in their specific context.

These observations warn against becoming attached to specific definitions of features like NIE and NDE (natural indirect and direct effects; p. 435, Equation 17.8) because these may not correspond to the causal actions a researcher wishes to investigate in their particular model. Indeed, Kline's definitions for these terms apply only when the model is structured exactly as in Figure 4, and only when specific features of the postulated intervention are assumed (remember the demanded 0–1 coding of  $X$ , the possibility of non-dichotomous  $X$  values, the arbitrary scale for  $\beta_0$ , and the required 0 for  $e_M$ ). Introducing additional model variables, and/or additional interactions or nonlinearities will change the equation for  $Y$  and/or the equations for the variables “replaced” in  $Y$ 's equation (as  $M$ 's equation replaced  $M$  in  $Y$ 's equation to obtain Equation 19 above). Such changes stymie any routine definition of entities such as NIE because they introduce terms not addressed by NIE.

Mean-centering  $M$  and  $Y$  (by subtracting the means from the appropriate data values) would set the intercept terms to zero, and hence eliminate some terms and alter the appropriate definitions, but would require countervailing un-centering if the interpretations were to be applied to variables having their original scales. Mean-centering an intervention like  $X$  would likely introduce confusion, because a 0 would no longer correspond to absence of the treatment, or 1 to presence of the treatment.



## Editorial Corrections and Other Small Improvements

- 1) Page xiii: the Chapter 6 and 7 titles should both contain or both omit the hyphen between Observed and Variables.
- 2) I fail to understand why Chapter 2 begins, opposite to usual traditions, by using upper-case B for unstandardized effects and lower-case b for standardized effects. This might be helpful if LISREL notation was to be used routinely, but that seems not to be the case.
- 3) The phrase “controlling for their intercorrelation” two lines above the equation on the middle of page 30 is inaccurate—with  $R^2$  there is no controlling.
- 4) Equation 2.12 is not a regression equation. The equation should include variables on the right side for it to be a regression equation.
- 5) The right sides of equations 2.12 and 2.13 are identical and cannot correspond to the different kinds of entities on the left of these equations.
- 6) Page 80, second line: should read “final 1–7” not 0–7.
- 7) Table 4.2 would be easier to read if the rescaled variances were presented in italics.
- 8) In both Equations 4.8 and 4.9, the term “ $r_{XX} \times r_{XX}$ ” should read “ $r_{XX} \times r_{YY}$ ”.
- 9) Page 133, first sentence: should be reworded to claim *only* no direct causal connection between  $X$  and  $Y$  because there is in fact some other causal connection between the variables.
- 10) Page 133, footnote 5: should end with “but this practice is not consistent with SEM”.
- 11) The discussion of causal loops would benefit from references clarifying how loops function (Hayduk 1987: Chapter 8) and how loops alter effects that touch any variable in a loop (Hayduk 1996: Chapter 3).
- 12) Page 151: The reference to “in the next chapter” at the end of the middle paragraph would be more helpful if it pointed specifically to “Instrumental Variables” (p. 180).
- 13) Page 161, last line of 1st paragraph: “excluded variables” should be “excluded effects”, and some additional headings could clarify the implicit segmentation of Appendix 7.A.
- 14) Page 167, last line: Instead of “a back-door path that starts” it would be more accurate to say, “a back-door path between  $X$  and  $Y$  that starts”.
- 15) Page 168, first line: Instead of “A back-door path may convey a spurious association between variables at either end, but never causation”, it would more accurate to say, “A back-door path may convey a spurious association between the variables at the ends of the path, but not a direct or indirect effect between the variables at the ends of the path.”
- 16) Page 168, second-last line of the 1st paragraph: The statement that the models “are equivalent” should instead clarify that the models are NOT causally equivalent (they contain contrasting causal effects) even though they imply equivalent-conditional-independencies.
- 17) Page 170, end of 1st paragraph: The claim that “multiple regression assumes no causal effects among the predictors” is incorrect. Multiple regression does not estimate/report effects among the predictors, but it does not assume these do not exist.
- 18) Page 170, paragraph 2, third-last line: Replace “where  $X$ , is specified to directly cause  $Y$ ” with “where  $X_1$  is specified to directly cause  $Y_1$ ”.
- 19) Page 175, end of second-last sentence in the 2nd paragraph: refers to  $X$  and  $Y$  but should refer to  $X_1$  and  $Y_1$ .
- 20) Page 177, 3rd paragraph, last line: should end “(see Appendix 8.A)”.
- 21) Page 180: The section on “Instrumental Variables” should clarify that the rules locate what the model claims as instrumental variables, but does not guarantee the corresponding worldly variables actually are instrumental variables (because the model may be misspecified). Farther on (page 182, paragraph 2), it is stated: “Exogenous variables make ideal instruments because by definition they are unrelated to all disturbances in the model.” The world is not controlled by this definition, and hence a researcher’s exogenous variables may or may not be unrelated to the modeled disturbances, and hence may or may not be acceptable instruments.
- 22) Page 189: “factor indeterminacy” should be “factor score indeterminacy”.
- 23) Page 197, 3rd paragraph, first line: “affect” should be “effect”.
- 24) Page 199, first line: Add the names of the Greek characters so this reads “( $\lambda$ , lambda)” and “( $\phi$ , phi)”.
- 25) Page 211: The term “matrix” which appears twice in the first two lines could be more helpfully described as a “list” or “vector”.

- 26) Page 212, bottom paragraph, line 6: Delete the word “indirectly” because the effects of latents on the indicators that measure those latents are direct.
- 27) Page 217, first line. The word “unreliability” should be replaced with “invalidity”.
- 28) Page 217, third last line in the first full paragraph: “improve factor measurement” should be “improve latent measurement” because the latent need not be a factor.
- 29) Page 223: 2nd paragraph, second line: “underlying factor” should read “underlying latent”.
- 30) Page 224: Figure 10.7 is referred to as Figure 11.7 in the website material.
- 31) Page 227: There seems no reason to switch the placement of the  $\xi$  and  $\eta$  terms away from their usual LISREL location in Appendix Equation 10.6, and this matrix equation is missing an  $\eta$  from the left side.
- 32) Page 241: The formula producing the equation on this page should be provided.
- 33) Page 241, 3rd paragraph, third line: should read “(see Figures 7.5 and 11.1)”.
- 34) Exercise 1 (pages 241 and 479) should incorporate the relevance of control/uncontrolled variables and the fact that this model fails significantly because both these features alter the permissible interpretations.
- 35) Page 261: An easy and generally applicable way to obtain start values for effects in complicated models can be obtained by making the square of the effect’s start-value multiplied by the variance of the causal variable contribute a reasonable amount of variance into the dependent variable—paralleling the structure of the terms in essay Equation 7 above. Checking that the square of an effect estimate multiplied by the variance of the cause contributes a reasonable amount of variance into the dependent variable can sometimes provide useful direction for finding where a model is empirically underidentified (p. 157).
- 36) Page 273 just prior to Equation 12.4: The word “limit” is statistically incorrect and should be deleted or replaced.
- 37) Page 275: The  $H_0$  near mid-page is missing a decimal, and should read “ $\geq .10$ ”.
- 38) Page 276: When readers encounter  $\hat{\Delta}_M$  just prior to Equation 12.7, they would probably appreciate a reminder that this was defined in Equation 12.4 (p. 273).
- 39) Page 307, line 2: should read “failed at  $p < .01$ ” because the model  $p = .006$  is larger than .001.
- 40) Page 358: Figure 14.4 is inconsistent with the website model output (two indicators are switched and the 1.0 connected to Risk is differentially placed).
- 41) Page 417, above the two equations: “Factor variances and sizes” should be “Factor variances and sample sizes”.
- 42) Page 438, second-last and third-last lines: should report how the feature is denoted, for example, as “the nonproduct factors A and B are all zero *which we denote as*  $\sigma_{AB,A}^2 = 0 = \sigma_{AB,B}^2$ ”.
- 43) Pages 438–39: Just as an introductory sentence indicates that the Equations in 17.11 were obtained “by taking the products of the corresponding expressions” there should have been an introductory sentence indicating how the Equations in 17.12 are obtained. The statement might indicate the first equation in the set of Equations 17.12 corresponds to the first equation in the set of equations ending column-1 in Kenny and Judd 1984:210; and might note that the remaining equations in Kline’s 17.12 parallel the first equation in the set with the last/covariance term repeatedly being dropped due to the assumed independence of error variables.
- 44) Page 439, the first line of text should read: where the term  $\sigma_{AB}^2$  *represents the variance of the product and the term*  $\sigma_{A,B}^2$  *represents the square of the covariance between factors A and B.*
- 45) Page 439: the third line of text should read “factors A and B plus *the square of their covariance*”.
- 46) Pages 452, 453, and 497: The year of the Hoyle and Isherwood reference should be 2013.
- 47) Page 455, last two lines: “have positive intercorrelations” should read “have substantial intercorrelations consistent with the signs of the indicators’ loadings on the latent factor”.
- 48) Page 456 first line: Delete “or negative”.
- 49) Page 457 in the last bullet point: “ensure your readers” should either be “assure your readers” or change the sentence to read “to ensure that it is actually”.
- 50) Page 497: Delete one of the duplicated journal titles for the Hoyle and Isherwood reference.
- 51) Page 507: The volume and pages for the Valeri and VanderWeele reference should read “18(2):137–50”.
- 52) Page 526: The entry for “Model test statistics” should NOT include “fit indexes” because the indices are not tests.