

Let's talk about Thurstone & Co.: An information-theoretical model for comparative judgments, and its statistical translation

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Abstract

This study revisits Thurstone's law of comparative judgment (CJ), focusing on two prominent issues of traditional approaches. First, it critiques the heavy reliance on Thurstone's Case V assumptions and, by extension, the Bradley-Terry-Luce (BTL) model when analyzing CJ data. Specifically, the study raises concerns about the assumptions of equal discriminial dispersions and zero correlation between the stimuli. While these assumptions simplify the trait measurement model, they may fail to capture the complexity of CJ data, potentially leading to unreliable and inaccurate trait estimates. Second, the study highlights the apparent disconnect between CJ's trait measurement and hypothesis testing processes. Although separating these processes simplifies the analysis of CJ data, it may also undermine the reliability of various statistical results derived from these processes.

To address these issues, the study extends Thurstone's general form using a systematic and integrated approach based on Causal and Bayesian inference methods. This extension integrates core theoretical principles alongside key assessment design features relevant to CJ experiments, such as the selection of judges, stimuli, and comparisons. It then translates these elements into a probabilistic statistical model for analyzing dichotomous CJ data, overcoming the rigid assumptions of Case V and the BTL model.

Finally, the study emphasizes the relevance of this extension for contemporary empirical CJ research, particularly stressing the need for bespoke CJ models tailored to the experiments and data assumptions. It also lays the foundation for broader applications, encouraging researchers across the social sciences to adopt more robust and interpretable methodologies.

Keywords: causal inference, directed acyclic graphs, structural causal models, bayesian statistical methods, thurstonian model, comparative judgement, probability, statistical modeling

1. Introduction

In *comparative judgment* (CJ) studies, judges assess a specific trait or attribute across different stimuli by performing pairwise comparisons (Thurstone, 1927b,a). Each comparison produces a dichotomous outcome, indicating which stimulus is perceived to have a higher trait level. For example, when assessing writing quality, judges compare pairs of written texts (the stimuli) to determine the relative writing quality each text exhibit (the trait) (Laming, 2004; Pollitt, 2012b; Whitehouse, 2012; van Daal et al., 2016; Lesterhuis, 2018a; Coertjens et al., 2017; Goossens and De Maeyer, 2018; Bouwer et al., 2023).

Numerous studies have documented the effectiveness of CJ in assessing traits and competencies over the past decade. These studies have highlighted three aspects of the method’s effectiveness: its reliability, validity, and practical applicability. Research on reliability suggests that CJ requires a relatively modest number of pairwise comparisons (Verhavert et al., 2019; Crompvoets et al., 2022) to generate trait scores that are as precise and consistent as those generated by other assessment methods (Coertjens et al., 2017; Goossens and De Maeyer, 2018; Bouwer et al., 2023). In addition, the evidence suggests that the reliability and time efficiency of CJ are comparable, if not superior, to those of other assessment methods when employing adaptive comparison algorithms (Pollitt, 2012b; Verhavert et al., 2022; Mikhailiuk et al., 2021). Meanwhile, research on validity indicates the capacity of CJ scores to represent accurately the traits under measurement (Whitehouse, 2012; van Daal et al., 2016; Lesterhuis, 2018a; Bartholomew et al., 2018; Bouwer et al., 2023). Lastly, research on its practical applicability highlights CJ’s versatility across both educational and non-educational contexts (Kimbell, 2012; Jones and Inglis, 2015; Bartholomew et al., 2018; Jones et al., 2019; Marshall et al., 2020; Bartholomew and Williams, 2020; Boonen et al., 2020).

Nevertheless, despite the increasing number of CJ studies, research in this domain remains unsystematic and fragmented, leaving several critical issues unresolved. This study identifies and discusses two prominent issues of traditional approaches that can undermine the reliability and validity of CJ’s trait estimates. First, it critiques the heavy reliance on Thurstone’s Case V assumptions (Thurstone, 1927a) and, by extension, the Bradley-Terry-Luce (BTL) model (Bradley and Terry, 1952; Luce, 1959) when analyzing CJ data. Specifically, the study raises concerns about the assumptions of equal discriminial dispersions and zero correlation between the stimuli. While these

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assumptions simplify the trait measurement model, they may fail to capture the complexity of CJ data, potentially leading to unreliable and inaccurate trait estimates. Second, the study highlights the disconnect between CJ’s trait measurement and hypothesis testing processes. Although separating these processes simplifies the analysis of CJ data, it may also undermine the reliability of various statistical inferences derived from these processes.

To address these issues, this study extends Thurstone’s general form through a systematic and integrated approach that combines Causal and Bayesian inference methods. In addition to potentially enhancing measurement reliability and validity, and improving statistical accuracy in hypothesis testing, this approach offers two key advantages. First, it clarifies the interactions among all actors and processes involved in CJ experiments. Second, it shifts the current comparative data analysis paradigm from passively accepting the BTL model assumptions to actively testing whether those assumptions fit the data under analysis.

As a result, the study divides its content into six main sections. Section 2 provides an overview of Thurstone’s theory. Section 3 discusses the identified issues in detail. Section 4 extends Thurstone’s general form to address these challenges. The extension integrates core theoretical principles alongside key CJ experimental design features, such as the selection of judges, stimuli, and comparisons. Section 5 translates these theoretical and practical elements into a probabilistic statistical model to analyze dichotomous pairwise comparison data. Finally, Section 6 discusses the findings, explores avenues for future research, and detail the challenges for future researchers.

2. Thurstone’s theory

In its most general form, Thurstone’s theory addresses pairwise comparisons wherein a single judge evaluates multiple stimuli (Thurstone, 1927a). The theory posits that two key factors determine the dichotomous outcome of these comparisons: the discriminial process of each stimulus and their discriminial difference. The *discriminal process* captures the psychological impact each stimulus exerts on the judge or, more simply, his perception of the stimulus trait. The theory assumes that the discriminial process for any given stimulus forms a Normal distribution along the trait continuum (Thurstone, 1927a). The mode (mean) of this distribution, known as the *modal discriminial process*, indicates the stimulus position on this continuum, while its dispersion, referred to as the *discriminal dispersion*, reflects variability in the perceived trait of the stimulus.

Figure 1a illustrates the hypothetical discriminial processes along a quality trait continuum for

two written texts. The figure indicates that the modal discriminative process for Text B is positioned further along the continuum than that of Text A ($T_B > T_A$), suggesting that Text B exhibits higher quality. Additionally, the figure highlights that Text B has a broader distribution compared to Text A, which arises from its larger discriminative dispersion ($\sigma_B > \sigma_A$).

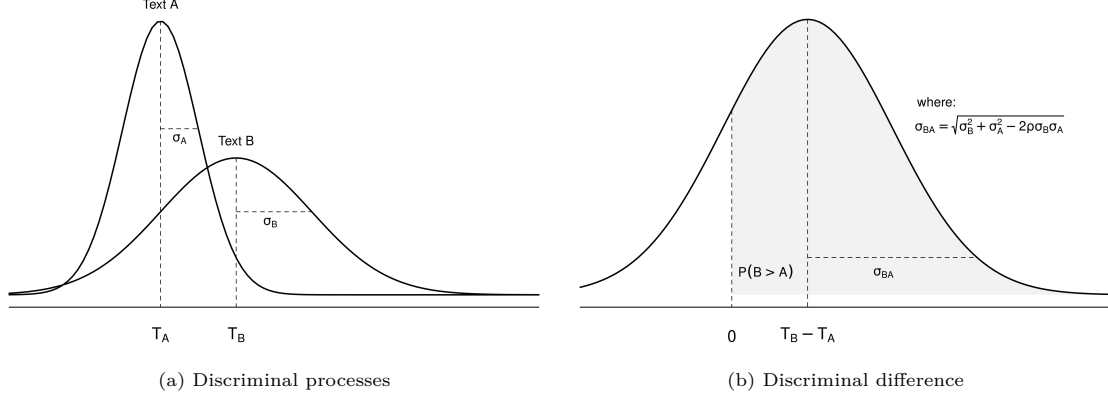


Figure 1: Hypothetical discriminative processes and discriminant difference along a quality trait continuum for two written texts.

However, since the individual discriminative processes of the stimuli are not directly observable, the theory introduces the *law of comparative judgment*. This law posits that in pairwise comparisons, a judge perceives the stimulus with a discriminative process positioned further along the trait continuum as possessing more of the trait (Bramley, 2008). This suggests that pairwise comparison outcomes depend on the relative distance between stimuli, not their absolute positions on the continuum. Indeed, the theory assumes that the difference between the underlying discriminative processes of the stimuli, referred to as the *discriminative difference*, determines the observed dichotomous outcome. Furthermore, the theory assumes that because the individual discriminative processes form a Normal distribution on the continuum, the discriminative difference will also conform to a Normal distribution (Andrich, 1978). In this distribution, the mode (mean) represents the average relative separation between the stimuli, and its dispersion indicates the variability of that separation.

Figure 1b illustrates the distribution of the discriminative difference for the two hypothetical texts. The figure indicates that the judge perceives Text B as having significantly higher quality than Text A. Two key observations support this conclusion: the positive difference between their modal discriminative processes ($T_B - T_A > 0$) and the probability area where the discriminative difference distinctly favors Text B over Text A, represented by the shaded gray area denoted as $P(B > A)$. As a result, the dichotomous outcome of this comparison is more likely to favor Text B over A.

Table 1: Thurstones cases and their assumptions

Assumption	General form	Thurstone's					BTL model
		Case I	Case II	Case III	Case IV	Case V	
Discriminal process (distribution)	Normal	Normal	Normal	Normal	Normal	Normal	Logistic
Discriminal dispersion (between stimuli)	Different	Different	Different	Different	Similar	Equal	Equal
Correlation (between stimuli)	One per pair	Constant	Constant	Zero	Zero	Zero	Zero
How many judges compare?	Single	Single	Multiple	Multiple	Multiple	Multiple	Multiple

3. Two Prominent Issues in Traditional CJ Practice

Thurstone noted from the outset that his general form, described in Section 2, led to a *trait scaling problem*. Specifically, the model required estimating more “unknown” parameters than the number of available pairwise comparisons (Thurstone, 1927a). For instance, in a CJ experiment with five texts, the general form would require estimating 20 parameters: five modal discriminative processes, five discriminative dispersions, and 10 correlations—one per comparison (see Table 1). However, a single judge could only provide $\binom{5}{2} = 10$ unique comparisons, an insufficient data set to estimate the required parameters.

To address this issue and facilitate the practical implementation of the theory, Thurstone developed five cases derived from this general form, each progressively incorporating additional simplifying assumptions (Thurstone, 1927a). In Case I, Thurstone postulated that pairs of stimuli would maintain a constant correlation across all comparisons. In Case II, he allowed multiple judges to undertake comparisons instead of confining evaluations to a single judge. In Case III, he posited that there was no correlation between stimuli. In Case IV, he assumed that the stimuli exhibited similar dispersions. Finally, in Case V, he replaced this assumption with the condition that stimuli had equal discriminative dispersions. Table 1 summarizes the assumptions of the general form and the five cases. For a detailed discussion of these cases and their progression, refer to Thurstone (1927a) and Bramley (2008, pp. 248–253).

However, Thurstone developed Case V prioritizing statistical simplicity over precise trait measurement and offering no guidance on how to use its trait estimates for statistical inference or hypothesis testing. Specifically, Thurstone cautioned that its use “should not be made without (an) experimental test” (Thurstone, 1927a, pp. 270), as it imposes the most extensive set of simplifying assumptions (Bramley, 2008; Kelly et al., 2022) (see Table 1). Moreover, because Thurstone’s primary goal was to produce a “rather coarse scaling” of traits and “allocate the compared stim-

uli on this continuum” (Thurstone, 1927a, pp. 269), his theory did not support formal statistical inference. Despite these limitations, it is surprising that CJ research has predominantly relied on Case V to measure different traits, which raises significant concerns about the reliability and validity of such measurements in contexts where the case’s assumptions may not hold (Kelly et al., 2022; Andrich, 1978). Furthermore, although the CJ tradition has attempted to address the gap of hypothesis testing by using the point estimates of the traits—or their transformations—a critical question remains: Does this approach provide a valid foundation for statistical inference?

Thus, this section discusses these two prominent issues. Specifically, Section 3.1 examines the heavy reliance on Thurstone’s Case V assumptions in the statistical analysis of CJ data. Conversely, Section 3.2 focuses on the apparent disconnect between the approaches to trait measurement and hypothesis testing in CJ.

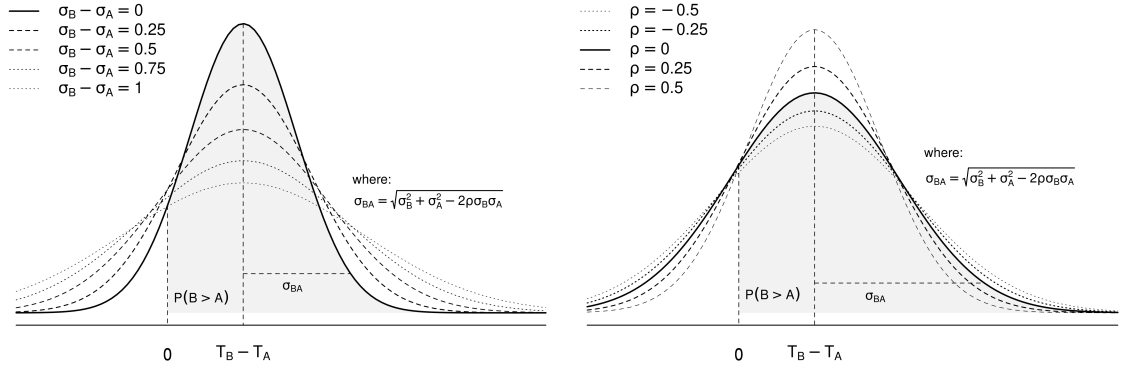
3.1. The Case V and the statistical analysis of CJ data

As previously discussed, Case V remains the most widely used model in CJ literature. This preference largely stems from the widespread adoption of the BTL model, which provides a simplified statistical representation of the case. The BTL model mirrors most of Case V’s assumptions, with one notable distinction: while Case V assumes a Normal distribution for the stimuli’ discriminial processes, the BTL model uses the more mathematically tractable Logistic distribution (Andrich, 1978; Bramley, 2008) (see Table 1). However, this substitution has minimal impact on trait estimation or model interpretation because the scale of the discriminial process (i.e., the latent trait) is arbitrary up to a non-monotonic transformation (van der Linden, 2017a; McElreath, 2021). That is, as long as the substitution (transformation) preserves the data rank order, the choice of distribution for the discriminial processes is inconsequential. This condition is satisfied in this case, as the Normal and Logistic distributions exhibit analogous statistical properties, differing only by a scaling factor of approximately 1.7 (van der Linden, 2017a).

However, Thurstone acknowledged that some assumptions of Case V could be problematic when researchers assess complex traits or heterogeneous stimuli (Thurstone, 1927b). Thus, given that modern CJ applications often involve such traits and stimuli, two key assumptions of Case V, and by extension, the BTL model, may not always hold in theory or practice. These assumptions are the equal dispersion and zero correlation between stimuli.

3.1.1. The assumption of equal dispersions between stimuli

According to the theory, discrepancies in the discriminial dispersions of stimuli shape the distribution of the discriminial difference, directly influencing the outcome of pairwise comparisons. A thought experiment can help illustrate this idea. In it, researchers observe the discriminial processes for two texts, A and B, assuming that the dispersion for Text A remains constant and that the two texts are uncorrelated ($\rho = 0$). Figure 2a demonstrates that an increase in the uncertainty associated with the perception of Text B relative to Text A ($\sigma_B - \sigma_A$), broadens the distribution of their discriminial difference. This broadening affects the probability area where the discriminial difference distinctly favors Text B over Text A, expressed as $P(B > A)$, ultimately influencing the comparison outcome. Additionally, the figure reveals that when the discriminial dispersions of the texts are equal, as in the BTL model ($\sigma_B - \sigma_A = 0$), the discriminial difference distribution is more narrow than when the dispersions differ. As a result, the discriminial difference is more likely to favor Text B over Text A, as it is represented by the shaded gray area.



(a) Discriminal Difference distribution under varying discrepancies in stimuli dispersions (b) Discriminal Difference distribution under varying levels of correlation between stimuli

Figure 2: The effect of dispersion discrepancies and stimuli correlation on the distribution of the discriminial difference.

In experimental practice, however, the thought experiment occurs in reverse. Researchers first observe the comparison outcome and then use the BTL model to infer the discriminial difference between stimuli and their respective discriminial processes (Thurstone, 1927b). Consequently, the outcome's ability to reflect *true* differences between stimuli largely depends on the validity of the model's assumptions (Kohler et al., 2019), in this case, the assumption of equal dispersions. For instance, when the assumption accurately captures the complexity of the data, the BTL model estimates a discriminial difference distribution that accurately represents the *true* discriminial difference between the texts. This scenario is illustrated in Figure 2a, when the model's discriminial difference distribution aligns with the *true* discriminial difference distribution, represented by the

thick continuous line corresponding to $\sigma_B - \sigma_A = 0$. The accuracy of this discriminial difference then ensures reliable estimates for the texts’ discriminial processes.

Notably, while assuming equal dispersions simplifies the trait measurement model, evidence from the CJ literature suggests that this assumption may fail to capture the complexity of modern CJ data. In particular, the assumption may not hold when researchers assess complex traits or heterogeneous stimuli (Thurstone, 1927b; Bramley, 2008; Kelly et al., 2022), as these traits and stimuli can introduce judgment discrepancies due to their unique characteristics (van Daal et al., 2016; Lesterhuis, 2018b; Chambers and Cunningham, 2022). Indeed, the CJ literature may already provide indications of such discrepancies, particularly in the form of “misfit” statistics. For instance, *misfit texts* are those whose comparisons result in more judgment discrepancies than those involving other texts (Pollitt, 2004, 2012b,a; Goossens and De Maeyer, 2018). These misfit texts may exhibit larger discriminial dispersions or represent outliers—texts with distinctive characteristics that deviate markedly from the rest of the sample (Grubbs, 1969). In either case, the BTL model neither accounts for these cases nor offers any means to address them, except their elimination from the analysis.

Significant statistical and measurement issues can arise when the assumption of equal dispersions between stimuli does not hold. Specifically, the BTL model may overestimate the trait’s reliability, that is, the degree to which the outcome accurately reflects the *true* discriminial differences between stimuli. This overestimation, in turn, results in spurious conclusions about these differences (McElreath, 2020) and, by extension, about the underlying discriminial processes of stimuli. Figure 2a also illustrates this scenario when the model’s discriminial difference distribution aligns with the thick continuous line for $\sigma_B - \sigma_A = 0$, while the *true* discriminial difference follows any discontinuous line where $\sigma_B - \sigma_A \neq 0$. Furthermore, if researchers acknowledge that misfit statistics may represent texts with different dispersions or outlying observations, the common CJ practice of excluding stimuli based on these statistics, as seen in Pollitt (2012a), Pollitt (2012b), van Daal et al. (2016), and Goossens and De Maeyer (2018), may unintentionally discard valuable information, introducing bias into the trait estimates (Zimmerman, 1994; McElreath, 2020). The direction and magnitude of these biases remain unpredictable, as they depend on which stimuli researchers exclude from the analysis.

3.1.2. The assumption of zero correlation between stimuli

The correlation ρ , illustrated in Figure 1b, measures how much the judges’ perception of a specific trait in one stimulus depends on their perception of the same trait in another. Similar to the

discriminal dispersions, this correlation shapes the distribution of the discriminial difference, directly impacting the outcomes of pairwise comparisons. A thought experiment, akin to the one presented in Section 3.1.1, can illustrate this idea. Assuming that the discriminial dispersions for both texts remain constant, Figure 2b shows that as the correlation between the two texts increases, the distribution of their discriminial difference becomes narrower. This narrowing, in turn, affects the probability that the discriminial difference distinctly favors Text B over Text A—denoted as $P(B > A)$ —and thus directly influences the comparison outcome. Furthermore, the figure shows that when two texts are independent or uncorrelated, as assumed in the BTL model ($\rho = 0$), the distribution of their discriminial difference is less narrow than in scenarios where the texts are positively correlated. As a result, it becomes less likely for the comparison to favor Text B over Text A, as indicated by the larger shaded area.

Despite these notable differences in the distribution of the discriminial difference under various correlational assumptions, in practice, experimental designs often adopt the assumption of no correlation between stimuli based on an old theoretical justification. Specifically, [Thurstone \(1927a\)](#) argued that stimuli could be treated as uncorrelated because judges’ biases—arising from two opposing and equally weighted effects occurring during the pairwise comparisons—would cancel each other out. This idea was later formalized by [Andrich \(1978\)](#), who provided a mathematical demonstration of this cancellation using the BTL model under the assumption of discriminial processes with additive biases. However, evidence from the CJ literature indicates that the assumption of zero correlation does not hold in practice in at least two scenarios: when intricate aspects of multidimensional, complex traits or heterogeneous stimuli influence judges’ perceptions or when additional hierarchical structures are relevant to the stimuli.

In the first scenario, research on text quality suggests that when judges evaluate multidimensional, complex traits or heterogeneous stimuli, they often rely on various intricate characteristics of the stimuli to form their judgments ([van Daal et al., 2016](#); [Lesterhuis, 2018b](#); [Chambers and Cunningham, 2022](#)). These additional relevant characteristics, when assessed, are unlikely to be equally weighted or opposing. As a result, they may exert an uneven influence on judges’ perceptions, creating biases that resist cancellation. For example, this could occur when a judge assessing the argumentative quality of a text places more weight on its grammatical accuracy than other judges, thereby favoring texts with fewer errors but weaker arguments. Furthermore, since the discriminial difference of the stimuli becomes an observable outcome only through the judges’ perceptions, these biases could introduce dependencies between the stimuli ([van der Linden, 2017b](#)). While direct ev-

idence for this particular scenario is lacking, studies such as [Pollitt and Elliott \(2003\)](#), [van Daal et al. \(2016\)](#) and [Bartholomew et al. \(2020\)](#) demonstrate the presence of such biases, supporting the notion that the factors influencing pairwise comparisons may not always cancel out.

In the second scenario, the shared context or inherent connections introduced by additional hierarchical structures may create dependencies between stimuli—a statistical phenomenon known as clustering ([Everitt and Skrondal, 2010](#)). For instance, when the same individual produces multiple texts, those texts often exhibit common features, such as writing style or overall quality, that judges can easily recognize. In this regard, although the CJ literature acknowledges the existence of such hierarchical structures, the statistical approaches to account for this additional source of dependence have been insufficient. For instance, when CJ data incorporates multiple samples of stimuli from the same individuals, researchers frequently rely on (averaged) point estimates of the BTL scores to conduct subsequent analyses and tests at the individual level ([Bramley and Vitello, 2019](#); [Boonen et al., 2020](#); [Bouwer et al., 2023](#); [van Daal et al., 2017](#); [Jones et al., 2019](#); [Gijzen et al., 2021](#)). However, this approach can introduce additional statistical and measurement issues, similar to the ones discussed in Section 3.2.

Thus, erroneously assuming zero correlation between stimuli can also lead to significant statistical and measurement issues. Specifically, neglecting judges’ biases or relevant hierarchical structures can create dimensional mismatches in the model, leading to the over- or underestimation of trait reliability ([Ackerman, 1989](#); [Hoyle, 2023](#)). These inaccuracies can result in spurious conclusions about the discriminial differences ([McElreath, 2020](#)) and, by extension, the underlying discriminial processes of the stimuli. This issue is illustrated in Figure 2b when the discriminial difference distribution of the BTL scores follows the thick continuous line ($\rho = 0$), while the *true* discriminial difference follows any discontinuous line where $\rho \neq 0$.

Finally, as discussed in the previous section, removing *misfit* judges—that is, judges whose assessments deviate markedly from the shared consensus ([Pollitt, 2012a,b](#); [van Daal et al., 2016](#); [Goossens and De Maeyer, 2018](#)), and considered outliers under the BTL model ([Grubbs, 1969](#))—risk discarding valuable information and introducing bias into trait estimates. The direction and magnitude of these biases remain unpredictable, as they depend on which judges researchers exclude from the analysis ([Zimmerman, 1994](#); [O’Hagan, 2018](#); [McElreath, 2020](#)).

3.2. The disconnect between trait measurement and hypothesis testing

Building on the previous section, it is clear that, researchers in CJ studies typically use the BTL model to measure traits and position the compared stimuli along a latent continuum (Thurstone, 1927a). The CJ literature also show that they frequently rely on point estimates of these traits (i.e., the BTL scores)—or their transformations—to conduct statistical inference or hypothesis testing. For example, researchers have used these scores to identify ‘misfit’ judges and stimuli (Pollitt, 2012b; van Daal et al., 2016; Goossens and De Maeyer, 2018), detect biases in judges’ ratings (Pollitt and Elliott, 2003; Pollitt, 2012b), calculate correlations with other assessment methods (Goossens and De Maeyer, 2018; Bouwer et al., 2023), or test hypotheses related to the underlying trait of interest (Casalicchio et al., 2015; Bramley and Vitello, 2019; Boonen et al., 2020; Bouwer et al., 2023; van Daal et al., 2017; Jones et al., 2019; Gijzen et al., 2021).

Nevertheless, while separating the trait measurement and hypothesis testing processes simplifies the analysis of CJ data, the statistical literature cautions against relying solely on the point estimates of BTL scores to conduct statistical inference or hypothesis tests, as this practice can undermine the resulting statistical conclusions. A key consideration is that BTL scores are parameter estimates that inherently carry uncertainty (measurement error). Ignoring this uncertainty can bias the analysis and reduce the precision of hypothesis tests. The direction and magnitude of such biases are often unpredictable. Results may be attenuated, exaggerated, or remain unaffected depending on the degree of uncertainty in the scores and the actual effects being tested (McElreath, 2020; Kline, 2023; Hoyle, 2023). Furthermore, the reduced precision in hypothesis tests diminishes their statistical power, increasing the likelihood of committing type-I or type-II errors (McElreath, 2020).

In aggregate, the heavy reliance on Thurstone’s Case V assumptions in the statistical analysis of comparative data can compromise the reliability of trait estimates. This overreliance may also undermine their validity (Perron and Gillespie, 2015), particularly when coupled with the disconnect between the trait measurement and hypothesis testing processes. However, the structural approach to causal inference can address these issues by offering a systematic and integrated framework that strengthens measurement reliability and validity while enhancing the statistical accuracy of hypothesis tests.

4. Extending Thurstone’s general form

The *structural approach* to causal inference provides a formal framework for identifying causes and estimating their effects using data. The approach uses structural causal models (SCMs) and directed acyclic graphs (DAGs) (Pearl, 2009; Pearl et al., 2016; Gross et al., 2018; Neal, 2020) to formally and graphically represent the assumed causal structure of a system, such as the one found in CJ experiments. Essentially, SCMs and DAGs function as *conceptual models* on which identification analysis rests. *Identification analysis* determines whether an estimator can accurately compute an estimand based solely on its (causal) assumptions, regardless of random variability (Schuessler and Selb, 2023). Here, *estimands* represent the specific quantities researchers aim to determine (Everitt and Skrondal, 2010). *Estimators* denote the methods or functions that transform data into an estimate, while *estimates* are the numerical values approximating the estimand (Neal, 2020; Everitt and Skrondal, 2010).

A motivating example that will appear in the rest of the document clarifies these concepts. In this example, researchers aim to determine: “To what extent do different teaching methods influence students’ ability to produce high-quality written texts?” To investigate this, a researcher designs a CJ experiment by randomly assigning students (individuals) to two groups, each receiving a different teaching method. Judges then compare pairs of students’ written texts (stimuli) to produce a dichotomous outcome reflecting the relative quality of each text (trait). Based on this setup, researchers can reformulate the research question as the estimand: “*On average*, is there a difference in the ability to produce high-quality written texts between the two groups of students?”. Following current CJ practices, researchers rely on estimates from the BTL model, or its transformations, to approximate this estimand.

However, Section 3 presents compelling evidence that Thurstone’s Case V, and by extension the BTL model, suffers from several statistical and measurement limitations. These limitations hinder the model’s ability to identify various estimands relevant to CJ inquiries, including the one described in the example. Identification is crucial because it is a necessary condition for ensuring consistent estimators. *Consistency* refers to the property of an estimator whose estimates converge to the “true” value of the estimand as the data size approaches infinity (Everitt and Skrondal, 2010). Without identification, consistency cannot be achieved, even with “infinite” and error-free data. Thus, deriving meaningful insights from finite data becomes impossible (Schuessler and Selb, 2023).

Fortunately, SCMs and DAGs support identification analysis through two key advantages¹. First, regardless of complexity, they can represent various causal structures using only five fundamental building blocks (Neal, 2020; McElreath, 2024). This feature allows researchers to decompose complex structures into manageable components, facilitating their analysis. Second, they depict causal relationships in a non-parametric way. This flexibility enables feasible identification strategies without requiring specification of the types of variables, the functional forms relating them, or the parameters of those functional forms (Pearl et al., 2016).

Thus, this section addresses the issues identified in Section 3 by extending Thurstone’s general form using the structural approach to Causal inference. Specifically, it combines the core theoretical principles outlined in Section 2 with key assessment design features relevant to CJ experiments, such as the selection of judges, stimuli, and comparisons. In addition to improving statistical accuracy and strengthening measurement reliability and validity, the approach offers two key advantages. First, it clarifies the interactions among all actors and processes involved in CJ experiments. Second, it shifts the current comparative data analysis paradigm from passively accepting the model assumptions to actively testing whether those assumptions fit the data under analysis.

Accordingly, Section 4.1 incorporates the theoretical principles into what we refer to as the *conceptual-population model*. This model assumes an idealized scenario where researchers have access to a *conceptual population* of comparative data, that is, data representing all repeated judgments made by every available judge for each pair of stimuli produced by each pair of individuals in the population. Conversely, Section 4.2 integrates the assessment design features into what we refer to as the *sample-comparison model*. This model assumes a more realistic scenario where researchers only have access to a sample of judges, individuals, stimuli, and comparisons from the conceptual population.

4.1. The conceptual-population model

In the conceptual-population model, the idealized scenario of a *conceptual population* of comparative data enables the integration of Thurstone’s theoretical principles and provides a foundation for proposing innovations aimed at addressing some of the issues discussed in Section 3.

¹These topics are beyond the scope of this study, thus, readers seeking a more profound understanding can refer to introductory papers such as Pearl (2010), Rohrer (2018), Pearl (2019), and Cinelli et al. (2020), and introductory books like Pearl and Mackenzie (2018), Neal (2020), and McElreath (2020) are useful. For more advanced study, seminal papers such as Neyman (1923), Rubin (1974), Spirtes et al. (1991), and Sekhon (2009), along with books such as Pearl (2009), Morgan and Winship (2014), and Hernán and Robins (2020), are recommended.

4.1.1. Integrating the first theoretical principles

Before incorporating the first theoretical principles of Thurstone’s theory, it is essential to further define SCMs. SCMs are formal mathematical models characterized by a set of *endogenous* variables V , a set of *exogenous* variables E , and a set of functions F (Pearl, 2009; Pearl et al., 2016; Cinelli et al., 2020). Endogenous variables are those whose causal mechanisms a researcher chooses to model (Neal, 2020). In contrast, exogenous variables represent *errors* or *disturbances* arising from omitted factors that the investigator chooses not to model explicitly (Pearl, 2009). Lastly, the functions, referred to as *structural equations*, express the endogenous variables as non-parametric functions of other endogenous and exogenous variables. These functions use the symbol ‘ $:=$ ’ to denote the asymmetrical causal dependence between variables and the symbol ‘ \perp ’ to represent *d-separation*, a concept akin to statistical (conditional) independence.

SCM 3a presents the first theoretical principles embedded in the conceptual-population model, which evaluates the impact of different teaching methods on students’ writing ability. This SCM outlines the relationship between the conceptual-population outcome (O_{iahbjk}^{cp}) and several related variables. The subscripts i and h identify the students who authored the texts (i.e., the individuals). The indices a and b represent the texts under comparison (i.e., the stimuli). The index j indicates the judge conducting the comparison, while the index k accounts for experimental conditions where a judge compares the same pair of stimuli multiple times, i.e., a *repeated measures designs* (Lawson, 2015, pp. 366-376). Thus, the indexing system supports comparisons between different texts written by the same student ($i = h; a \neq b$) and between texts written by distinct students ($i \neq h$; where $a = b$ is permitted), each compared once or repeatedly by all judges ($j = 1, \dots, n_J; k = 1, \dots, n_K$; where $n_J > 1$ and $n_K \geq 1$). However, it excludes cases where a judge compares a student’s text to itself, whether once or multiple times ($i = h; a = b; j = 1, \dots, n_J; k = 1, \dots, n_K$; where $n_J > 1$ and $n_K \geq 1$), as such comparison lacks practical relevance within the CJ framework. Here, n_J indicates the total number of judges, and n_K denotes the number of repeated judgments each judge performs.

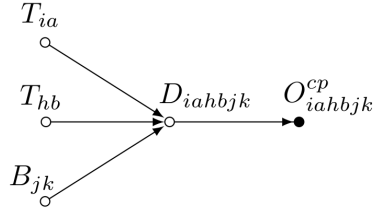
In line with Thurstone’s theory, SCM 3a depicts the texts’ discriminial processes (T_{ia}, T_{hb}) and their discriminial difference (D_{iahbjk}) (see Section 2). Additionally, the SCM incorporates a key design feature of CJ experiments: the judges’ biases (B_{kj}). This extension builds on the arguments presented in Section 3.1.2, contending that the discriminial difference becomes an observable outcome only through judges’ perceptions. Given that such perceptions may be imperfect—and that each judge may carry some degree of bias (see Pollitt and Elliott, 2003; van Daal et al., 2016)—it is

reasonable that judges' perceptions (bias) should be treated as an integral component of the CJ system from the outset, as this leads to a more accurate representation of the data-generating process underlying the pairwise comparisons. This model defines the preliminary set of endogenous variables, $V = \{O_{iahbjk}, D_{iahbjk}, T_{ia}, T_{hb}, B_{kj}\}$, and the preliminary set of structural equations, $F = f_O, f_D$, which capture the non-parametric dependencies among these variables.

$$O_{iahbjk}^{cp} := f_O(D_{iahbjk})$$

$$D_{iahbjk} := f_D(T_{ia}, T_{hb}, B_{kj})$$

(a) SCM



(b) DAG

Figure 3: Conceptual-population model, scalar form.

Notably, every SCM has an associated DAG (Pearl et al., 2016; Cinelli et al., 2020). A DAG is a *graph* consisting of nodes connected by edges, where nodes represent random variables. The term *directed* indicates that edges or arrows extend from one node to another, indicating the direction of causal influence. The absence of an edge implies no direct relationship between the nodes. The term *acyclic* means that the causal influences do not form loops, ensuring the influences do not cycle back on themselves (McElreath, 2020). DAGs conventionally depict observed variables as solid black circles and unobserved (latent) variables as open circles (Morgan and Winship, 2014). Although DAGs conventionally omit exogenous variables for simplicity, the DAGs presented in this section includes exogenous variables to improve clarity and reveal potential issues related to conditioning and confounding (Cinelli et al., 2020).

Figure 3b displays the DAG corresponding to SCM 3a, illustrating the expected causal relationships outlined in Thurstone's theory. The graph shows that the discriminative processes of the texts (T_{ia}, T_{hb}) influence their discriminative difference (D_{iahbjk}), which in turn determines the outcome (O_{iahbjk}^{cp}). It also highlights the influence of judges' biases (B_{kj}) on the discriminative difference. Additionally, the DAG differentiates between observed endogenous variables, such as the outcome

(solid black circle), and latent endogenous variables, including the texts' discriminial processes, their discriminial difference, and the judges' biases (open circles).

4.1.2. The conceptual-population data structure

Although specifying a data structure is not mandatory when using SCMs and DAGs, defining one in this case can improve clarity and facilitate the description of the system. Thus, to re-express the scalar form of the CJ system shown in Figure 3 into an equivalent vectorized form, we first define the vectors I and J , along with the matrices IA and JK , as follows:

$$I = \begin{bmatrix} 1 \\ \vdots \\ i \\ \vdots \\ h \\ \vdots \\ n_I \end{bmatrix} ; J = \begin{bmatrix} 1 \\ \vdots \\ j \\ \vdots \\ n_J \end{bmatrix} ; IA = \begin{bmatrix} 1 & 1 \\ \vdots & \vdots \\ 1 & n_A \\ \vdots & \vdots \\ i & a \\ \vdots & \vdots \\ h & b \\ \vdots & \vdots \\ n_I & 1 \\ \vdots & \vdots \\ n_I & n_A \end{bmatrix} ; JK = \begin{bmatrix} 1 & 1 \\ \vdots & \vdots \\ 1 & n_K \\ \vdots & \vdots \\ j & k \\ \vdots & \vdots \\ n_J & 1 \\ \vdots & \vdots \\ n_J & n_K \end{bmatrix} \quad (1)$$

Here, each element of I represents a unique individual i or h , where n_I denotes the total number of individuals. Similarly, each element of J corresponds to a unique judge j , with n_J indicating the total number of judges. Moreover, each row of IA represents a unique pairing of individuals i, h with stimuli a, b . As a result, the matrix IA contains $n_I \cdot n_A$ rows and 2 columns, where n_A specifies the number of stimuli available per individual. Likewise, each row of JK associates a judge j with a (repeated) judgment index k . Consequently, the matrix JK has $n_J \cdot n_K$ rows and 2 columns, where n_K indicates the number of repeated judgments each judge makes.

Additionally, we construct the matrix R to map each row of the IA matrix with a corresponding row from the JK matrix. This matrix has n rows and 6 columns, where $n = \binom{n_I \cdot n_A}{2} \cdot n_J \cdot n_K$. Here, the term $\binom{n_I \cdot n_A}{2}$ represents the binomial coefficient, which quantifies the total number of unique comparisons possible between every pair of stimuli generated by each pair of individuals in the population. Thus, we define the matrix as follows:

$$R = \begin{bmatrix} 1 & 1 & 1 & 2 & 1 & 1 \\ \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\ 1 & 1 & 1 & 2 & 1 & n_K \\ \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\ i & a & h & b & j & k \\ \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\ n_I & n_A - 1 & n_I & n_A & n_J & 1 \\ \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\ n_I & n_A - 1 & n_I & n_A & n_J & n_K \end{bmatrix} \quad (2)$$

It is easier to visualize the structure of these vectors and matrices by considering an example. Assuming $n_I = 5$, $n_A = 2$, $n_J = 3$, and $n_K = 3$, the vectors and matrices described in Equations (1) and (2) take the following form:

$$I = \begin{bmatrix} 1 \\ 2 \\ 3 \\ 4 \\ 5 \end{bmatrix} ; J = \begin{bmatrix} 1 \\ 2 \\ 3 \end{bmatrix} ; IA = \begin{bmatrix} 1 & 1 \\ 1 & 2 \\ 2 & 1 \\ 2 & 2 \\ 3 & 1 \\ 3 & 2 \\ 4 & 1 \\ 4 & 2 \\ 5 & 1 \\ 5 & 2 \end{bmatrix} ; JK = \begin{bmatrix} 1 & 1 \\ 1 & 2 \\ 1 & 3 \\ 2 & 1 \\ 2 & 2 \\ 2 & 3 \\ 3 & 1 \\ 3 & 2 \\ 3 & 3 \end{bmatrix} ; R = \begin{bmatrix} 1 & 1 & 1 & 2 & 1 & 1 \\ 1 & 1 & 1 & 2 & 1 & 2 \\ 1 & 1 & 1 & 2 & 1 & 3 \\ \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\ 1 & 1 & 5 & 2 & 1 & 1 \\ 1 & 1 & 5 & 2 & 1 & 2 \\ 1 & 1 & 5 & 2 & 1 & 3 \\ \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\ 4 & 2 & 5 & 2 & 3 & 1 \\ 4 & 2 & 5 & 2 & 3 & 2 \\ 4 & 2 & 5 & 2 & 3 & 3 \\ 5 & 1 & 5 & 2 & 3 & 1 \\ 5 & 1 & 5 & 2 & 3 & 2 \\ 5 & 1 & 5 & 2 & 3 & 3 \end{bmatrix} \quad (3)$$

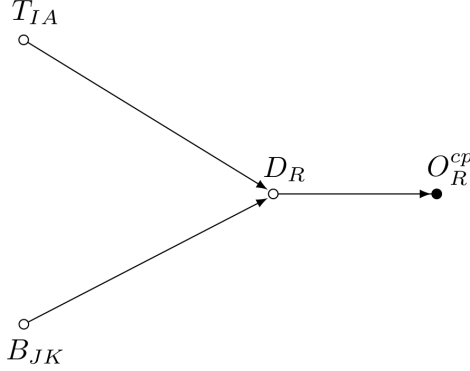
Now, using Equations (1) and (2), we can re-express SCM 3a and DAG 3b in an equivalent vectorized form, as shown in Figure 4. In this depiction, the outcome O_R^{cp} , the texts' discriminial difference D_R , their discriminial processes T_{IA} , and the judges' biases B_{JK} are represented as vectors rather than scalar values. These vectors capture all the observations from the conceptual population. Specifically, O_R^{cp} and D_R are observed and latent vectors of length n , respectively. Moreover, T_{IA}

and B_{JK} are latent vectors of lengths $n_I \cdot n_A$ and $n_J \cdot n_K$, respectively.

$$O_R^{cp} := f_O(D_R)$$

$$D_R := f_D(T_{IA}, B_{JK})$$

(a) SCM



(b) DAG

Figure 4: Conceptual-population model, initial vectorized form.

4.1.3. Integrating hierarchical structural components

Building on the principles of Structural Equation Modeling (SEM) (Hoyle, 2023) and Item Response Theory (IRT) (Fox, 2010; van der Linden, 2017a), the conceptual-population model integrates two *hierarchical structural components* to examine how different teaching methods influence students’ writing ability. Each structural component defines how observed or latent variables affect the primary latent variable of interest (Everitt and Skrandal, 2010). The model’s hierarchical design allows researchers to formulate and test hypotheses that account for both the nested structure of stimuli and the uncertainties inherent in trait estimation (see Section 3.1.2 and Section 3.2 for a discussion of these considerations).

The top branch of DAG 5b illustrates the first component, where *relevant*² student-related variables X_I , such as teaching method, and students’ idiosyncratic errors e_I causally influence the latent variable representing students’ writing-quality trait T_I . The error term e_I captures variations in

²*Relevant variables* are those that satisfy the *backdoor criterion* (Neal, 2020, pp 37), that is, they belong to a *sufficient adjustment set* (Pearl, 2009; Pearl et al., 2016; Morgan and Winship, 2014). A *sufficient set* (potentially empty) blocks all non-causal paths between a predictor and an outcome without opening new ones (Pearl, 2009). Refer also to footnote 1.

students' traits unexplained by X_I . Here, X_I is an observed matrix with n_I rows and q_I independent columns (variables), and both e_I and T_I are latent vectors of length n_I . Additionally, this branch shows how T_I , along with *relevant*³ text-related variables X_{IA} (e.g., text length), and texts' idiosyncratic errors e_{IA} causally influence the texts' written-quality trait T_{IA} , the first primary latent variable of interest. The error term e_{IA} captures variations in the texts' traits that remain unexplained by T_I or X_{IA} . Here, X_{IA} is an observed matrix with dimensions $n_I \cdot n_A$ rows and q_{IA} independent columns (variables), while e_{IA} and T_{IA} are latent matrices with n_I rows and n_A columns.

Similarly, the bottom branch of DAG 5b depicts the second component, where *relevant*⁴ judge-related variables Z_J , such as judgment expertise, and judges' idiosyncratic errors e_J causally influence the latent variable representing judges' bias B_J . The error e_J captures variations in judges' bias unexplained by Z_J . Here, Z_J is an observed matrix with n_J rows and q_J independent columns (variables), and both e_J and B_J are latent vectors of length n_J . Furthermore, the branch shows how B_J , along with *relevant*⁵ judgment-related variables Z_{JK} (e.g., the number of judgments a judge makes), and judgments' idiosyncratic errors e_{JK} causally influence the judges' biases associated with each text B_{JK} , the second primary latent variable of interest. The error e_{JK} captures variations in judgments unexplained by B_J or Z_{JK} . Here, Z_{JK} is an observed matrix with dimension $n_J \cdot n_K$ rows and q_{JK} independent columns (variables), while e_{JK} and B_{JK} are latent matrices with n_J rows and n_K columns.

Notably, all variables and functions shown in SCM 5a and DAG 5b are part of the set of endogenous variables V , structural equations F , and exogenous variables E for the conceptual-population model. Additionally, the figures demonstrate that all exogenous variables are independent of one another, as indicated by the relationships $e_{IA} \perp \{e_I, e_{JK}, e_J\}$, $e_I \perp \{e_{JK}, e_J\}$ and $e_{JK} \perp e_J$ and the absence of connecting arrows.

Overall, the conceptual-population model extends Thurstone's general form by introducing key innovations to address the limitations discussed in Section 3.1.2 and Section 3.2. These enhancements include accounting for judges' biases and integrating hierarchical structural components. Nevertheless, despite its promise of enhancing measurement accuracy and precision, the model still depends on the unrealistic assumption that researchers have access to a *conceptual population*. Since

³refer to footnote 2.

⁴refer to footnote 2.

⁵refer to footnote 2.

researchers rarely meet this assumption in practice, they must consider a more realistic scenario.

$$O_R^{cp} := f_O(D_R)$$

$$D_R := f_D(T_{IA}, B_{JK})$$

$$T_{IA} := f_T(T_I, X_{IA}, e_{IA})$$

$$T_I := f_T(X_I, e_I)$$

$$B_{JK} := f_B(B_J, Z_{JK}, e_{JK})$$

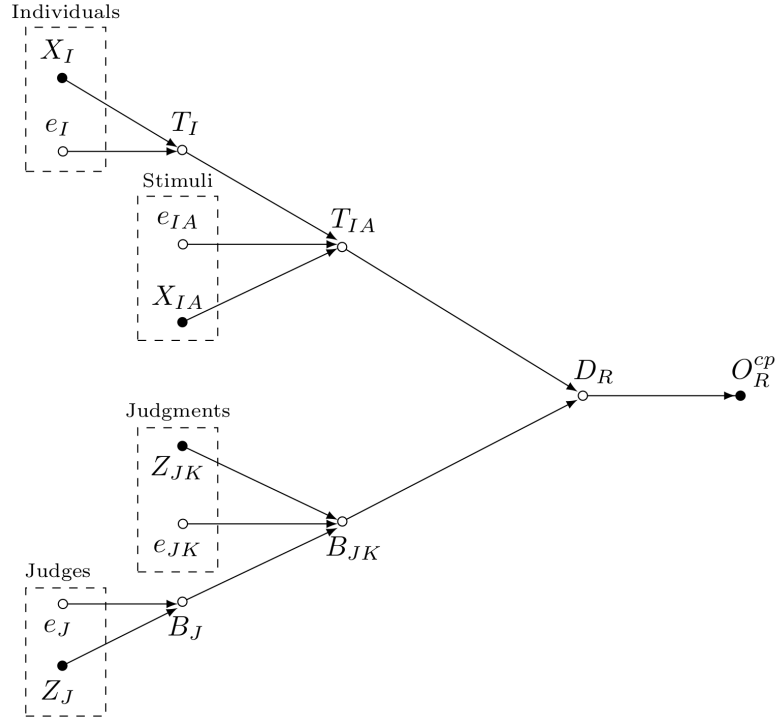
$$B_J := f_B(Z_J, e_J)$$

$$e_I \perp \{e_J, e_{IA}, e_{JK}\}$$

$$e_J \perp \{e_{IA}, e_{JK}\}$$

$$e_{IA} \perp e_{JK}$$

(a) SCM



(b) DAG

Figure 5: Conceptual-population model, final vectorized form.

4.2. The sample-comparison model

The sample-comparison model presents a more realistic scenario than the conceptual-population model. First, in Section 4.2.1, it explicitly assumes researchers work with a data sample consisting of a limited number of repeated judgments (n_K^s) from a sample of judges (n_J^s) and a specific number of texts (n_A^s) from a sample of students (n_I^s), all drawn from the conceptual population. Second, in Section 4.2.2, the model assumes that judges do not perform *all repeated judgments* within the data sample. Instead, they conduct a sufficient number of stimuli comparisons, n_C , to ensure an accurate estimation of the proportion $P(B > A)$, as proposed by [Thurstone \(1927a\)](#).

4.2.1. The sample mechanism

To incorporate the sampling mechanism and facilitate the interpretation of the sample-comparison model, we first define the *data sampling process* using the binary vector variables S_I , S_J , S_{IA} , and S_{JK} as follows:

$$S_I = \begin{bmatrix} i_{(1)} \\ \vdots \\ i_{(i)} \\ \vdots \\ i_{(h)} \\ \vdots \\ i_{(nI)} \end{bmatrix} ; S_J = \begin{bmatrix} j_{(1)} \\ \vdots \\ j_{(j)} \\ \vdots \\ j_{(nJ)} \end{bmatrix} ; S_{IA} = \begin{bmatrix} ia_{(1,1)} \\ \vdots \\ ia_{(1,n_A)} \\ \vdots \\ ia_{(i,a)} \\ \vdots \\ ia_{(h,b)} \\ \vdots \\ ia_{(nI,1)} \\ \vdots \\ ia_{(nI,n_A)} \end{bmatrix} ; S_{JK} = \begin{bmatrix} jk_{(1,1)} \\ \vdots \\ jk_{(1,n_K)} \\ \vdots \\ jk_{(j,k)} \\ \vdots \\ jk_{(nJ,1)} \\ \vdots \\ jk_{(nJ,nK)} \end{bmatrix} \quad (4)$$

Where each element of S_I is a binary value indicating the presence or absence of corresponding elements in the vector I , as in Equation (5). We apply the same logic to S_J using vector J (not shown). Thus, the vectors S_I and S_J contains n_I and n_J elements, respectively.

$$i_{(i)} = \begin{cases} 1 & \text{if data element } i \text{ from } I \text{ is sampled} \\ 0 & \text{if data element } i \text{ from } I \text{ is missing} \end{cases} \quad (5)$$

Similarly, each element of S_{IA} is a binary value indicating the presence or absence of data rows in the matrices IA , as defined in Equation (6). We apply the same logic to S_{JK} using the matrix JK

(not shown). Thus, the vectors S_{IA} and S_{JK} contains $n_I \cdot n_A$ and $n_J \cdot n_K$ elements, respectively.

$$ia_{(i,a)} = \begin{cases} 1 & \text{if data elements } i, a \text{ from } IA \text{ are sampled} \\ 0 & \text{if data elements } i, a \text{ from } IA \text{ are missing} \end{cases} \quad (6)$$

We can visualize these vectors more clearly using the example in Equation (3). Suppose researchers exclude the second student and the third judge, and they also omit the second text from each individual and the third repeated judgment from each judge. Given $n_I = 5$, $n_A = 2$, $n_J = 3$, and $n_K = 3$, these vectors take the following structure:

$$S_I = \begin{bmatrix} 1 \\ 0 \\ 1 \\ 1 \\ 1 \end{bmatrix} ; S_J = \begin{bmatrix} 1 \\ 1 \\ 0 \end{bmatrix} ; S_{IA} = \begin{bmatrix} 1 \\ 0 \\ 0 \\ 1 \\ 0 \\ 1 \\ 0 \\ 1 \\ 0 \end{bmatrix} ; S_{JK} = \begin{bmatrix} 1 \\ 1 \\ 0 \\ 1 \\ 1 \\ 0 \\ 0 \\ 0 \\ 0 \end{bmatrix} \quad (7)$$

Equation (7) shows that missing observations in the vectors S_I and S_J —which represent unsampled students and judges—directly determine which observations are missing in S_{IA} and S_{JK} . In other words, researchers can only observe texts and judgments from students and judges initially included in the sample. The equation also shows that the sum of observed elements in S_I equals the number of sampled students (n_I^s) and that a similar sum in vector S_J equals the sampled judges (n_J^s). Conversely, the sum of observed elements in S_{IA} represents the total sampled texts across all sampled students ($n_I^s \cdot n_A^s$), while a similar sum in vector S_{JK} represents the total sampled repeated judgments across all sampled judges ($n_J^s \cdot n_K^s$). Notice that in this example, because the design systematically excludes every third repeated judgment, researchers can also express S_{JK} using $n_K = n_K^s = 2$.

Finally, we define the *sample mechanism* S in Equation (8), which maps each element of S_{IA} to every element of S_{JK} . Each element $s_{(i,a,h,b,j,k)}$ is a binary value indicating the presence or absence of data rows in the matrix R resulting from the sample mechanism, as in Equation (9). Thus,

the vector contains n elements, matching the number of rows in R , and the sum of its elements represents the total data sample: $n^s = \binom{n_I^s \cdot n_A^s}{2} \cdot n_J^s \cdot n_K^s$. Here, the term $\binom{n_I^s \cdot n_A^s}{2}$ represents the binomial coefficient, which quantifies the total number of unique comparisons possible between every pair of sampled stimuli generated by each pair of sampled individuals.

$$S = \begin{bmatrix} s_{(1,1,1,2,1,1)} \\ \vdots \\ s_{(1,1,1,2,1,n_K)} \\ \vdots \\ s_{(i,a,h,b,j,k)} \\ \vdots \\ s_{(n_I,n_A-1,n_I,n_A,n_J,1)} \\ \vdots \\ s_{(n_I,n_A-1,n_I,n_A,n_J,1)} \end{bmatrix} \quad (8)$$

$$s_{(i,a,h,b,j,k)} = \begin{cases} 1 & \text{if data elements } i, a, h, b, j, k \text{ from } R \text{ are sampled} \\ 0 & \text{if data elements } h, i, a, b, j, k \text{ from } R \text{ are missing} \end{cases} \quad (9)$$

With the definition of S , we incorporate the sample mechanism into the conceptual-population model. Following the convention of [McElreath \(2020\)](#) and [Deffner et al. \(2022\)](#), DAG 6b represents the conceptual-population outcome O_R^{cp} as unobserved, emphasizing that researchers cannot directly access this outcome due to the sampling mechanism. The DAG also depicts the *sample design* vector S as a causal factor influencing the sample-comparison outcome O_R^{sc} . A square encloses this vector, indicating that it is a conditioned variable. Here, *conditioning* means that researchers restrict their focus to the elements of O_R^{cp} that satisfy $s_{(i,a,h,b,j,k)} = 1$ ([Neal, 2020](#); [McElreath, 2020](#)). Essentially, S serves as the vector selecting *all repeated judgments a subset of judges makes for the subset of stimuli produced by the sampled individuals*.

Notably, the DAG shows that S is independent of all other variables in the model. This implies that DAG 6b applies exclusively to Simple Random Sampling (SRSg) designs. In these designs, each repeated judgment, judge, stimulus, and individual has the same probability of being included in the sample as any other observation within their respective groups ([Lawson, 2015](#)).

$$O_R := f_C(O_R^{sc}, C)$$

$$O_R^{sc} := f_S(O_R^{cp}, S)$$

$$O_R^{cp} := f_O(D_R)$$

$$D_R := f_D(T_{IA}, B_{JK})$$

$$T_{IA} := f_T(T_I, X_{IA}, e_{IA})$$

$$T_I := f_T(X_I, e_I)$$

$$B_{JK} := f_B(B_J, Z_{JK}, e_{JK})$$

$$B_J := f_B(Z_J, e_J)$$

$$e_I \perp \{e_J, e_{IA}, e_{JK}\}$$

$$e_J \perp \{e_{IA}, e_{JK}\}$$

$$e_{IA} \perp e_{JK}$$

(a) SCM



(b) DAG

Figure 6: Sample-comparison model, final vectorized form

However, due to concerns about the practical feasibility of the comparison task ([Boonen et al.](#),

2020), CJ experiments rarely implement an exhaustive pairings of sampled judges, stimuli, and individuals. Thus, a realistic scenario must account for the fact that judges typically compare only a subset of stimuli authored by certain individuals.

4.2.2. The comparison mechanism

As in the previous section, we begin defining the *comparison mechanism* using the binary vector variable C to facilitate the interpretation of the sample-comparison model. Equation (10) shows that C contains n elements corresponding to the number of rows in the R matrix, with each element $c_{(i,a,h,b,j,k)}$ being a binary value indicating the presence or absence of data rows in R , a definition similar to that of $s_{(i,a,h,b,j,k)}$ in Equation (9).

$$C = \begin{bmatrix} c_{(1,1,1,2,1,1)} \\ \vdots \\ c_{(1,1,1,2,1,n_K)} \\ \vdots \\ c_{(i,a,h,b,j,k)} \\ \vdots \\ c_{(n_I,n_A-1,n_I,n_A,n_J,1)} \\ \vdots \\ c_{(n_I,n_A-1,n_I,n_A,n_J,1)} \end{bmatrix} \quad (10)$$

The DAG 6b also integrates the *comparison mechanism* C into the conceptual-population model. It shows the sample-comparison outcome as unobserved, highlighting that researchers cannot directly access to O_R^{sc} due to the comparison mechanism. Moreover, the DAG depicts C as a causal factor influencing O_R , enclosing it in a square to indicate that it is a conditioned variable. Specifically, C determines *which repeated judgments the subset of judges makes for the subset of stimuli produced by the sampled individuals*. In essence, C reflects the assumption that judges *do not* perform all possible repeated judgments for the selected stimuli. Instead, they carry out a “sufficient number of comparisons” n_C to allow the accurate estimation of the proportion $P(B > A)$ for each stimulus pair (Thurstone, 1927a, pp. 267).

Notably, DAG 6b also shows that C is independent of all other variables in the model. This independence implies that the conceptual model represented by the DAG applies exclusively to Random Allocation Comparative Designs (Bramley, 2015), or Incomplete Block Designs (Lawson, 2015), where every repeated judgment has an equal probability of being included in the sample.

$$O_R := f_O(D_R, S, C)$$

$$D_R := f_D(T_{IA}, B_{JK})$$

$$T_{IA} := f_T(T_I, X_{IA}, e_{IA})$$

$$T_I := f_T(X_I, e_I)$$

$$B_{JK} := f_B(B_J, Z_{JK}, e_{JK})$$

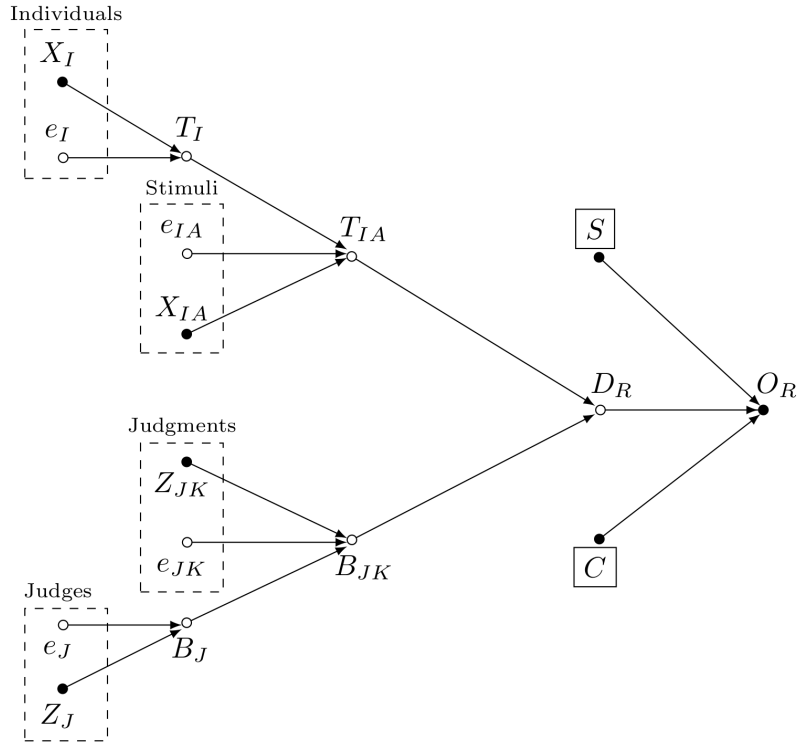
$$B_J := f_B(Z_J, e_J)$$

$$e_I \perp \{e_J, e_{IA}, e_{JK}\}$$

$$e_J \perp \{e_{IA}, e_{JK}\}$$

$$e_{IA} \perp e_{JK}$$

(a) SCM



(b) DAG

Figure 7: Comparative judgment model

Finally, since it is standard to assume that the distribution of the conceptual-population outcome O_R^{cp} also holds for O_R^{sc} and O_R , we can reformulate the sample-comparison model in Figure 6 into the equivalent form shown in Figure 7. This reformulation produces a model that applies directly to

a sample of comparative data. In this version, the unobserved outcomes O_R^{cp} and O_R^{sc} are omitted, and O_R inherits the structural equation f_O that originally defined O_R^{cp} . Moreover, the definition of O_R now reflects its direct dependence on the discriminial difference D_R and the sample and comparison mechanisms, S and C .

In summary, the SCM 7a and DAG 7b extend Thurstone’s general form to address several limitations of the BTL model. These extensions account for judge biases (see Section 4.1.1), reflect the hierarchical structure of stimuli and incorporate measurement error in trait estimation and hypothesis testing (see Section 4.1.3), and clarify the role of the sample and comparison mechanisms in CJ experiments (see Section 4.2). However, they do not resolve concerns about the assumption of equal dispersions among stimuli discussed in Section 3.1.1. Since this concern relates to the statistical assumption underlying the distribution of the discriminial process, we develop a formal statistical model to address it in the next section.

5. From SCM to statistical model

With the structure in SCM 7a, we can derive a statistical model that addresses violations of the equal dispersion assumption (see Section 3.1.1). This derivation is possible because a fully specified SCM encodes functional and probabilistic information, which we can replace with suitable functions and probabilistic assumptions (Pearl et al., 2016). Specifically, SCM 7a allows us to express the joint distribution of our complex CJ system as a product of simpler conditional probability distributions (CPDs)⁶, as shown in Equation (11). For clarity, we treat expressions such as $Y := f_Y(X)$, $P(Y | X)$, and $Y \sim f(Y | X)$ as equivalent, where $P(Y | X)$ and $f(Y | X)$ represent the CPD of Y given X .

$$\begin{aligned}
& P(O_R, S, C, D_R, T_{IA}, X_{IA}, e_{IA}, T_I, X_I, e_I, B_{JK}, Z_{JK}, e_{JK}, B_J, Z_J, e_J) \\
&= P(O_R | D_R, S, C) \cdot P(S) \cdot P(C) \cdot P(D_R | T_{IA}, B_{JK}) \\
&\quad \cdot P(T_{IA} | T_I, X_{IA}, e_{IA}) \cdot P(T_I | X_I, e_I) \\
&\quad \cdot P(B_{JK} | B_J, Z_{JK}, e_{JK}) \cdot P(B_J | Z_J, e_J) \\
&\quad \cdot P(X_{IA}) \cdot P(X_I) \cdot P(Z_{JK}) \cdot P(Z_J) \\
&\quad \cdot P(e_{IA}) \cdot P(e_I) \cdot P(e_{JK}) \cdot P(e_J)
\end{aligned} \tag{11}$$

Each CPD in Equation (11) rests on specific assumptions, which we outline in Figure 8. We begin

⁶This re-expression is possible because the *chain rule* of probability and the *Bayesian Network Factorization (BNF)* property. For further details, see Pearl et al. (2016) and Neal (2020).

by assuming that O_R follows a Bernoulli distribution⁷, reflecting the binary nature of CJ outcomes. Furthermore, following the conventions of Generalized Linear Models (GLMs) (McCullagh and Nelder, 1983; Lee and Nelder, 1996; Agresti, 2015), the distribution links O_R to the latent discriminial difference vector D_R using an inverse-logit function: $\text{inv_logit}(x) = 1/(1 + \exp(-x))$.

$O_R := f_O(D_R, S, C)$	$P(O_R D_R, S, C)$	$O_R \stackrel{iid}{\sim} \text{Bernoulli}[\text{inv_logit}(D_R)]$
$D_R := f_D(T_{IA}, B_{JK})$	$P(D_R T_{IA}, B_{JK})$	$D_R = (T_{IA}[i, a] - T_{IA}[h, b]) + B_{JK}[j, k]$
$T_{IA} := f_T(T_I, X_{IA}, e_{IA})$	$P(T_{IA} T_I, X_{IA}, e_{IA})$	$T_{IA} = T_I + \beta_{XA}X_{IA} + e_{IA}$
$T_I := f_T(X_I, e_I)$	$P(T_I X_I, e_I)$	$T_I = \beta_{XI}X_I + e_I$
$B_{JK} := f_B(B_J, Z_{JK}, e_{JK})$	$P(B_{JK} B_J, Z_{JK}, e_{JK})$	$B_{JK} = B_J + \beta_{ZK}Z_{JK} + e_{JK}$
$B_J := f_B(Z_J, e_J)$	$P(B_J Z_J, e_J)$	$B_J = \beta_{ZJ}Z_J + e_J$
$e_I \perp \{e_J, e_{IA}, e_{JK}\}$	$P(e_I)P(e_{IA})P(e_J)P(e_{JK})$	$e \sim \text{Multi-Normal}(\mu, \Sigma)$
$e_J \perp \{e_{IA}, e_{JK}\}$		$\Sigma = DRD$
$e_{IA} \perp e_{JK}$		
(a) SCM	(b) Probabilistic model	(c) Statistical model

Figure 8: Comparative judgment model, SCM, probabilistic and statistical model assuming different discriminial dispersions for the student’s traits

Note that while the joint distribution in Equation (11) includes the probability distributions of the sampling and comparison mechanisms, $P(S)$ and $P(C)$, as well as those of the predictor variables— $P(X_{IA})$, $P(X_I)$, $P(Z_{JK})$, and $P(Z_J)$ —we omit these probabilities from the models presented in Figure 8. We make this omission because, although these distributions contribute to the overall joint distribution of the data, the variables S , C , X_{IA} , X_I , Z_{JK} , and Z_J are observed and independent of any other variable in the model. As observed variables, they do not require distributional assumptions—unlike the idiosyncratic errors. Moreover, their independence stems from the random selection designs that govern the variables⁸.

⁷The binomial distribution—including its special case, the Bernoulli distribution—represent a maximum entropy distribution for binary events (McElreath, 2020, pp. 34). This means that the Bernoulli distribution is the most consistent alternative when only two un-ordered outcomes are possible and their expected frequencies are assumed to be constant (McElreath, 2020, pp. 310). For a detailed discussion of the binomial as a maximum entropy distribution, see McElreath (2020, sec. 10.1.2).

⁸Randomization ensures that data—and, by extension, an estimator—satisfies several key identification properties,

Next we define D_R as the difference between the discriminial processes $T_{IA}[i, a]$ and $T_{IA}[h, b]$, representing the underlying written-quality trait of the compared texts, plus the corresponding repeated judge bias $B_{JK}[j, k]$. Note that if we assume that $B_{JK}[j, k]$ reflects the difference in stimulus-specific biases, i.e., $B_{JK}[j, k] = B_{JK}[i, a, j, k] - B_{JK}[h, b, j, k]$, we can re-write the discriminial difference as:

$$\begin{aligned} D_R &= (T_{IA}[i, a] - T_{IA}[h, b]) + B_{JK}[j, k] \\ &= (T_{IA}[i, a] + B_{JK}[i, a, j, k]) - (T_{IA}[h, b] + B_{JK}[h, b, j, k]) \\ &= T_{IA}^*[i, a] - T_{IA}^*[h, b] \end{aligned} \tag{12}$$

This formulation reveals that the discriminial difference captures a *pure interaction effect*, in which neither the texts' discriminial processes nor the judges' biases alone determine the outcome, but their interaction does (Attia et al., 2022). Put simply, this mathematical description captures the idea that the stimuli' discriminial processes become an observable outcome only through the lens of judges' perceptions (i.e., their biases). For clarity, the square brackets in D_R indicate the relevant indices for each trait vector; they do not imply any subsetting of the data.

We now specify the functional forms for T_{IA} , T_I , B_{JK} , and B_J . We model T_{IA} as a linear combination of the students' underlying writing-quality traits T_I , the effects of relevant text-related variables on quality assessment $\beta_{XA}X_{IA}$ (such as the influence of text length), and the text-specific idiosyncratic errors e_{IA} . Similarly, we express T_I as a linear combination of relevant student-related variables affecting the quality assessment $\beta_{XI}X_I$, and student-specific idiosyncratic errors e_I . For the judge-specific terms, we model B_{JK} as a linear combination of the judge's individual bias B_J , the influence of relevant judgment-related variables on quality assessment $\beta_{ZK}Z_{JK}$ (e.g., how the number of judgments affect the evaluation), and judgment-specific idiosyncratic errors e_{JK} . Finally, we define B_J as a linear combination of relevant judge-level variables influencing the quality assessment $\beta_{ZJ}Z_J$ (such as judgment expertise) and judge-specific idiosyncratic errors e_J .

Next, we specify the probabilistic assumptions for the idiosyncratic errors e_I , e_{IA} , e_J , and e_{JK} . Unlike other variables in the model, these error terms exhibit indeterminacies in their *location*,

such as common support, no interference, and consistency. The most critical property, however, is the elimination of confounding. *Confounding* occurs when an external variable, such as X_I , simultaneously influences both the outcome (e.g., O_R) and a variable of interest (e.g., S), resulting in spurious associations between the latter two (Everitt and Skrondal, 2010). Randomization ensure the absence of confounding by effectively decoupling the association between the variable of interest and any other variable, except for the outcome itself. For a more detailed discussion on the benefits of randomization, see Pearl (2009), Morgan and Winship (2014), Neal (2020), and Hernán and Robins (2020).

orientation, and *scale* due to the lack of an inherent scale in the associated latent variables T_I , T_{IA} , B_J , and B_{JK} . Thus, to identify the latent variable model, we must resolve these indeterminacies (Depaoli, 2021; de Ayala, 2009). Drawing on principles from SEM (Hoyle, 2023), we assume that the vector of idiosyncratic errors $e = [e_I, e_{IA}, e_J, e_{JK}]^T$, follows a Multivariate Normal distribution with mean vector μ and a covariance matrix $\Sigma = DRD$, with D denoting a diagonal matrix of standard deviations and R a correlation matrix. To address the *location* indeterminacy, we set the errors' mean vector to zero:

$$\mu = [0, 0, 0, 0]^T \quad (13)$$

Following SCM 8a, we resolve the *orientation* indeterminacy by assuming that the errors are uncorrelated. This assumption leads us to specify a correlation matrix for the errors, R , as:

$$R = \begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{bmatrix} \quad (14)$$

To resolve the *scale* indeterminacy, we define the diagonal matrix D as:

$$D = \begin{bmatrix} s_{XI} & 0 & 0 & 0 \\ 0 & p_{IA} & 0 & 0 \\ 0 & 0 & s_{ZJ} & 0 \\ 0 & 0 & 0 & p_{JK} \end{bmatrix} \quad (15)$$

Here, s_{XI} represents the standard deviation for the individuals, p_{IA} for the stimuli, s_{ZJ} for the judges, and p_{JK} for the judgments. We assume s_{XI} varies depending on the teaching method group to which each student belongs. Using the example from Section 4, where the teaching method $X_I = \{1, 2\}$, the model sets the constraint according to Equation (16). This constraint anchors the scale of the individuals' latent trait while relaxing the assumption of equal dispersion for the stimuli, thereby addressing the concerns raised in Section 3.1.1.

$$\sum_{g=1}^2 s_{XI}[g]/2 = 1 \quad (16)$$

Note that because the error vector e follows an uncorrelated Multivariate Normal distribution, the marginal distribution of e_{IA} is a univariate Normal distribution with mean zero and standard deviation p_{IA} . We set P_{IA} as a proportion of 1 to establish the scale of the stimuli' latent trait

relative to the scale of the individuals' trait. As a result, T_{IA} is also normally distributed. This configuration effectively reinstates Thurstone's original assumption of Normal discriminial processes for the stimuli (see Table 1).

Similarly, we assume s_{ZJ} varies depending on the groups to which each judge belongs. For instance, if $Z_J = \{1, 2, 3\}$ represents three groups of judges with varying expertise, the model sets the constraint according to Equation (17). This constraint anchors the scale of the judges' latent trait and relaxes the assumption of equal dispersion for the judgments.

$$\sum_{g=1}^2 s_{ZJ}[g]/3 = 1 \quad (17)$$

Conversely, p_{JK} is defined as a proportion of 1 to establish the scale of the judgments' latent trait relative to the scale of the judges' trait.

Finally, we use *Bayesian inference methods* to convert model 8c into a practical statistical tool for analyzing paired comparison data. Bayesian inference offers three key advantages for this purpose. First, it handles complex and overparameterized models, where the number of parameters exceeds the number of observations (Baker, 1998; Kim and Cohen, 1999)—such as the model we try to implement. Second, it incorporates prior information to constrain parameter estimates within plausible bounds, thereby mitigating estimation issues like non-convergence or improper solutions that often affect frequentist methods (Martin and McDonald, 1975; Seaman III et al., 2011). In our implementation, we use priors to define the error distribution, which sets the scale of the latent variables (Depaoli, 2014). Third, Bayesian inference supports robust inferences from small samples, where the asymptotic properties underlying frequentist methods are less reliable (Baldwin and Fellingham, 2013; Lambert et al., 2006; Depaoli, 2014). This feature is particularly relevant in CJ experiments, where researchers often collect large volumes of paired comparisons but work with relatively small samples of judges, stimuli, and individuals to test hypotheses.

The **Declarations** section of this document provides a link to the model code, along with an alternative specification that assumes equal discriminial dispersions. We tested both versions of the model with success using **Stan** (Stan Development Team., 2021, version 2.26.1).

6. Discussion

Thurstone introduced the Law of Comparative Judgment to measure psychological traits of stimuli through pairwise comparisons (Thurstone, 1927b,a). In its general form, the theory models single-

judge comparisons across multiple, potentially correlated stimuli. Each comparison produces a dichotomous outcome indicating which stimulus the judge perceives as having a higher trait level. However, Thurstone identified one key challenge with its measurement model: it required estimating more “unknown” parameters than the number of available pairwise comparisons (Thurstone, 1927a). To address this issue and facilitate the theory’s practical applicability, he formulated five cases derived from this general form, each progressively incorporating several simplifying assumptions.

Of these five cases, the CJ literature has predominantly relied on Case V to measure various psychological traits, as shown in studies like Kimbell (2012), Jones and Inglis (2015), and Boonen et al. (2020). This preference largely stems from the widespread adoption of the BTL model, which offers a simplified statistical formulation of the case. The BTL model mirrors the assumptions of Case V—namely, equal discriminial dispersions and zero correlation for the stimuli’ discriminial processes—with one notable distinction: while Case V assumes normally distributed discriminial processes, the BTL model uses the more mathematically tractable Logistic distribution (Andrich, 1978; Bramley, 2008).

Although replacing the Normal with the Logistic distribution has a minimal impact on trait estimation or model interpretation (van der Linden, 2017a; McElreath, 2021), Thurstone acknowledged that the two remaining assumptions of Case V—and, by extension, of the BTL model—could pose significant problems when researchers assess complex traits or heterogeneous stimuli (Thurstone, 1927a). In particular, he noted that while assuming equal discriminial dispersions and zero correlation for discriminial processes simplifies the trait measurement model, these assumptions may fail to reflect the complexity of CJ data (Thurstone, 1927b; Andrich, 1978; Kelly et al., 2022). As a result, they can lead to unreliable and inaccurate trait estimates (Ackerman, 1989; Zimmerman, 1994; McElreath, 2020; Hoyle, 2023).

Furthermore, because Thurstone aimed primarily to produce a trait’s “coarse scaling” and “allocate the compared stimuli on this continuum” (Thurstone, 1927b, pp. 269), his theory did not specify how to use the resulting trait estimates for statistical inference. The CJ tradition has attempted to address this gap by separating trait estimation from the hypothesis testing process. Specifically, CJ studies often rely on point estimates of the traits (i.e., the BTL scores)—or their transformations—to conduct statistical inference. Although separating these processes simplifies the analysis of CJ data, the statistical literature cautions against this practice, as it can introduce bias into the analysis and compromise the reliability of the hypothesis test (McElreath, 2020; Kline,

2023; Hoyle, 2023).

Thus, to address the limitations of Thurstone’s Case V and the BTL model—particularly their strong assumptions and the disconnect between trait measurement and hypothesis testing—this study extends Thurstone’s general form using a systematic, integrated approach that combines Causal and Bayesian inference methods. The approach begins by formulating a conceptual model represented by an SCM and a DAG. This model integrates Thurstone’s core theoretical principles—such as the discriminial processes (i.e., traits) of stimuli—alongside key CJ experimental design features, including judges’ bias, sampling procedures, and comparison mechanisms.

The study then translates the SCM into a bespoke statistical model that addresses key limitations of Case V. This model allows researchers to analyze CJ data when the assumptions of equal dispersion and zero correlation do not hold (see Section 3.1.1 and Section 3.1.2), and when they need to perform statistical inference (refer to Section 3.2). In particular, the model accounts for judge biases (see Section 4.1.1), captures the hierarchical structure of stimuli (refer to Section 4.1.3), incorporates measurement error in the hypothesis testing process (refer to Section 4.1.3), and accommodates heterogeneity in discriminial dispersions (see Section 5). These methodological innovations can potentially enhance the reliability and validity of trait measurement (Perron and Gillespie, 2015) and improve the accuracy of statistical inferences.

Beyond these potential benefits, the approach offers two additional advantages. First, it clarifies the interactions among all actors and processes involved in CJ experiments. Second, it shifts the analytic paradigm from passively accepting the assumptions of the BTL model to actively testing their fit with observed data. Together, these advantages create a principled framework for evaluating best practices in CJ experimental designs, one that better aligns with the demands of contemporary CJ assessment contexts (Kelly et al., 2022) and opens new avenues for research.

Five research avenues deserve particular attention due to their direct impact on the reliability and validity of trait estimates and the accuracy of statistical inferences. First, researchers should investigate how sampling and comparison mechanisms influence CJ experimental outcomes and affect inference robustness. Second, studies should assess the impact of judges’ bias on the number of comparisons needed to achieve reliable trait estimates. Third, future work should re-evaluate current methods for identifying ‘misfitting’ judges and stimuli. Fourth, researchers should analyze how the hierarchical structure of stimuli affects both experimental design and hypothesis testing. Finally, studies should explore how modeling non-dichotomous outcomes or incorporating multidimensional

traits affects the reliability and validity of the trait estimates.

The explicit integration of the sampling and comparison mechanisms within our conceptual model offers a novel perspective on the first research avenue. Specifically, it shows how these mechanisms influence the data-generating process by acting as missing data sources, determining which observations are part of the data sample. This new perspective invites the application of Little and Rubin’s principled missing data framework (Little and Rubin, 2020), enabling a more rigorous evaluation of existing claims about these mechanisms, their influence on CJ outcomes, and their implications for designing and evaluating more complex experimental setups.

One prominent claim in the CJ literature is that Thurstone’s model naturally handles non-random missing data without compromising the reliability or validity of trait estimates (Bramley, 2008). However, it is easy to imagine scenarios where this assumption may not hold. For instance, selecting a particular group of judges could result in an incomplete representation of the trait’s full range and complexity, as judgment accuracy varies with demographic factors (Bartholomew et al., 2020; Kelly et al., 2022) and even between individuals (Gill and Bramley, 2013; van Daal et al., 2016, 2017; van Daal, 2020). Consequently, judges’ selection could affect the formation of a “shared consensus”, potentially biasing trait estimates and distorting the observed trait distribution (Deffner et al., 2022). It is therefore essential to determine whether such sampling biases occur—and if so, whether they undermine the validity of trait estimates and the robustness of statistical inferences. Our framework offers a rigorous setting to test the scenario.

another example: For instance, consider a CJ experiment assessing speech quality using a case-control design (Everitt and Skrondal, 2010) that samples individuals with normal hearing (NH) and those with hearing impairment (HI). Here, the sampling mechanism depends on hearing status—a variable likely to influence the trait of interest, as HI individuals may exhibit lower speech quality (Boonen et al., 2023; Rivera Espejo et al.). Suppose a subgroup within the HI population, such as individuals with genetically induced hearing loss, is difficult to reach. If this group is systematically excluded and happens to over- or under-perform in speech quality, the resulting trait estimates may be biased. It is therefore essential to determine whether such biases occur—and if so, whether they undermine the validity of trait estimates and the robustness of statistical inferences—and our framework provides a rigorous setting to test this claim.

another example: For instance, consider a high-stakes non-mandatory summative CJ assessment, where the results the individuals have in this assessment have some implications for them, however,

their participation is not mandatory. In this case, it not hard to imagine that some of these individuals will not participate in the CJ assessment, thus affecting the sample, and ultimately the trait estimates: now a probable under performing group will not available in the data sample, thus biasing the resulting trait estimates. It is therefore essential to determine whether such biases occur—and if so, whether they undermine the validity of trait estimates and the robustness of statistical inferences—and our framework provides a rigorous setting to test this claim.

Moreover, when designing and evaluating more complex experimental setups, our framework highlights one key principle: the sampling and comparison mechanisms directly influence the CJ outcome. In this study, we deliberately structured these mechanisms to produce data that is *missing completely at random* (MCAR)—that is, the missingness is unrelated to any observed or unobserved variables, including the outcome (Little and Rubin, 2020). This design offers one key advantage: it generates simple random samples that satisfy the condition of ignorability, enabling researchers to legitimately ignore missing data during analysis without introducing bias (Everitt and Skrondal, 2010; Kohler et al., 2019).

However, many modern CJ applications involve more complex forms of missingness—such as *missing at random* (MAR) or *missing not at random* (MNAR) mechanism (Little and Rubin, 2020)—that arise from the chosen sampling and comparison methods. Two notable examples are adaptive comparative judgment designs (Pollitt, 2012a,b) and case-control studies (Everitt and Skrondal, 2010). In adaptive designs, an algorithm defines the comparison mechanism by dynamically selecting which stimulus pairs judges evaluate. These designs have produced mixed evidence: some studies report improved trait reliability (Pollitt and Elliott, 2003; Pollitt, 2012a,b), while others suggest these gains may be artificially inflated (Bramley, 2015; Bramley and Vitello, 2019; Cromptoets et al., 2020, 2022). In contrast, case-control studies rely on sampling mechanisms linked to non-assignable variables that likely influence the trait of interest. For instance, Boonen et al. (2020) selected children based on hearing status, a non-assignable variable that likely affects speech quality (Boonen et al., 2023; Rivera Espejo et al.). Nevertheless, regardless of their missingness mechanism, both designs could benefit from explicitly articulating their underlying causal structures. Doing so would enable researchers to assess and test their impact on trait estimation and statistical inference. Our framework provides a rigorous foundation for such articulations.

Additionally, by explicitly incorporating judges’ biases, our framework provides a novel approach to investigate the second research avenue. Specifically, it enables researchers to determine whether

such biases occur and, if so, to assess their impact on the reliability and validity of trait estimation. Although some anecdotal evidence already suggests judges’ biases exist (Pollitt and Elliott, 2003; van Daal et al., 2016; Bartholomew et al., 2020), further research is needed to address several key questions. For instance: Can CJ data be analyzed under the assumption of “sample-free” trait calibration—that is, the estimated trait difference for any two scripts does not depend on which other scripts they are compared with, implying an absence of judges’ bias? (Bramley, 2008; Andrich, 1978) If this assumption does not hold, how many judges and judgments are required to ensure acceptable levels of trait reliability and validity? Do training and expertise influence judgment biases, and to what extent? (Kelly et al., 2022) Finally, what are the benefits of *repeated measures designs*, in which judges evaluate the same stimulus pairs multiple times? (Lawson, 2015)

About removing misfit judges and stimuli

About the hierarchical structure of stimuli

About non-dichotomous outcomes and multidimensional traits

While a natural step would be to formally derive answers to all these research avenues—similar to approaches used in classical models like linear regression—we argue that the complexity of the CJ system makes this route impractical. Instead, simulation-based methods, like power analysis, offer a more feasible alternative, allowing researchers to investigate these issues without relying on cumbersome mathematical proofs.

Despite its potential, this study presents several challenges that researchers must navigate. First, researchers need to be proficient in Probabilistic Programming Languages (PPLs) and Bayesian inference methods. This expertise is essential for translating context-specific conceptual models into bespoke statistical models that meaningfully address the research questions at hand. While such translations are not exclusive to Bayesian methods, we argue that Bayesian approaches offer a more accessible path. They reduce, for instance, the need for additional expertise in optimization techniques and theory often required by frequentist methods, thereby lowering the technical barrier for many applied researchers.

Second, although initial simulation tests of the statistical model have yielded promising results (not shown), these findings may not generalize across research contexts. We have argued that the model’s ability to accurately estimate traits and support valid statistical inferences depends heavily on the specific research context—particularly on whether the data and inferential goals align with

the model’s underlying assumptions. That said, our approach provides tools for researchers to test and evaluate these context-dependent CJ models directly.

Third,

Fourth, applying this approach requires a solid understanding of Causal inference. For instance, throughout the study, we assume that predictor variables must be “relevant” to the research context, interpreting relevance as the inclusion of the predictor variables in a “sufficient adjustment set.” However, we do not fully explore what this entails or its implications for model specification and validity. Nonetheless, we provide key references throughout the study to help motivated researchers explore these foundational concepts in greater depth.

Fifth and final, our approach does not establish a clear epistemological connection between the theoretical latent trait and its quantitative estimate. In other words, the approach does not provide a definitive answer to the fundamental question: How do we know that the estimated trait reflects the theoretical trait of interest? However, this limitation is not unique to our approach; it reflects a fundamental challenge all research involving latent constructs faces. Nevertheless, we argue that our approach offers methodological tools that enable researchers to design CJ experiments under conditions that support reliable and valid trait measurement.

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7. Appendix

7.1. Statistical and Causal inference

This section introduces fundamental statistical and causal inference concepts necessary for understanding the core theoretical principles described in this document. It does not, however, offer a comprehensive overview of causal inference methods. Readers seeking more in-depth understanding may wish to explore introductory papers such as [Pearl \(2010\)](#), [Rohrer \(2018\)](#), [Pearl \(2019\)](#), and [Cinelli et al. \(2020\)](#). They may also find it helpful to consult introductory books like [Pearl and Mackenzie \(2018\)](#), [Neal \(2020\)](#), and [McElreath \(2020\)](#). For more advanced study, readers may refer to seminal intermediate papers such as [Neyman \(1923\)](#), [Rubin \(1974\)](#), [Spirtes et al. \(1991\)](#), and [Sekhon \(2009\)](#), as well as books such as [Pearl \(2009\)](#), [Morgan and Winship \(2014\)](#), and [Hernán and Robins \(2020\)](#).

7.1.1. Empirical research and randomized experiments

Empirical research uses evidence from observation and experimentation to address real-world challenges. In this context, researchers typically formulate their research questions as *estimands* or *targets of inference*, i.e., the specific quantities they seek to determine ([Everitt and Skrondal, 2010](#)). For instance, researchers might be interested in answering the following question: “To what extent do different teaching methods (T) influence students’ ability to produce high-quality written texts (Y)?” To investigate this, researchers could randomly assign students to two groups, each exposed to a different teaching method ($T_i = \{1, 2\}$). Then, they would perform pairwise comparisons, generating a dichotomous outcome ($Y_i = \{0, 1\}$) showing which student exhibits more of the ability. In this scenario, the research question can be rephrased as the estimand, “*On average*, is there a difference in the ability to produce high-quality written texts between the two groups of students?” and this estimand can be mathematically represented by the random associational quantity in Equation 18, where $E[\cdot]$ denotes the expected value.

$$E[Y_i | T_i = 1] - E[Y_i | T_i = 2] \tag{18}$$

Researchers then proceed to identify the estimands. *Identification* determines whether an estimator can accurately compute the estimand based solely on its assumptions, regardless of random variability ([Schuessler and Selb, 2023](#), pp. 4). An *estimator* refers to a method or function that transforms data into an estimate ([Neal, 2020](#)). *Estimates* are numerical values that approximate

the estimand derived through the process of *estimation*, which integrates data with an estimator (Everitt and Skrondal, 2010). The Identification-Estimation flowchart (McElreath, 2020; Neal, 2020), shown in Figure 9, visually represents the transition from estimands to estimates.

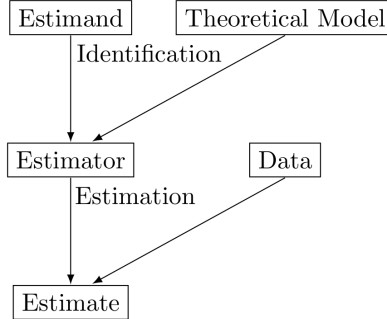


Figure 9: Identification-Estimation flowchart. Extracted and slightly modified from Neal (2020, pp. 32)

Identification is a necessary condition to ensure *consistent* estimators. An estimator achieves *consistency* when it converges to the “true” value of an estimand as the data size approaches infinity (Everitt and Skrondal, 2010). Without identification, researchers cannot achieve consistency, even with “infinite” and error-free data. As a result, deriving meaningful insights about an estimand from finite data becomes impossible (Schuessler and Selb, 2023, pp. 5). Therefore, to ensure accurate and reliable estimates, researchers prioritize estimators with desirable identification properties. For instance, the Z-test is a widely used estimator for comparing group proportions, yielding accurate estimates when its underlying assumptions are satisfied (Kanji, 2006). Furthermore, researchers can interpret estimates from the Z-test as causal, provided the data is collected through a randomized experiment.

Randomized experiments are widely recognized as the gold standard in evidence-based science (Hariton and Locascio, 2018; Hansson, 2014). This recognition stems from their ability to enable researchers interpret associational estimates as causal. They achieve this by ensuring data, and by extension an estimator, satisfies several key identification properties, such as common support, no interference, and consistency (Morgan and Winship, 2014; Neal, 2020). The most critical property, however, is the elimination of confounding. *Confounding* occurs when an external variable X simultaneously influences the outcome Y and the variable of interest T , resulting in spurious associations (Everitt and Skrondal, 2010). Randomization addresses this issue by decoupling the association between the intervention allocation T from any other variable X (Morgan and Winship, 2014; Neal, 2020).

Nevertheless, researchers often face constraints that limit their ability to conduct randomized experiments. These constraints include ethical concerns, such as the assignment of individuals to potentially harmful interventions, and practical limitations, such as the infeasibility of, for example, assigning individuals to genetic modifications or physical impairments (Neal, 2020). In these cases, causal inference offers a valuable alternative for generating causal estimates and understanding the mechanisms underlying specific data. In addition, the framework can provide significant theoretical insights that can enhance the design of experimental and observational studies (McElreath, 2020).

7.1.2. Identification under causal inference

Unlike classical statistical modeling, which focuses primarily on summarizing data and inferring associations, the *causal inference* framework is designed to identify causes and estimate their effects using data (Shaughnessy et al., 2010; Neal, 2020). The framework uses rigorous mathematical techniques to address the *fundamental problem of causality* (Pearl, 2009; Pearl et al., 2016; Morgan and Winship, 2014). This problem revolves around the question, “What would have happened ‘in the world’ under different circumstances?” This question introduces the concept of counterfactuals, which are instrumental in defining and identifying causal effects.

Counterfactuals are hypothetical scenarios that are *contrary to fact*, where alternative outcomes resulting from a given cause are neither observed nor observable (Neal, 2020; Counterfactual, 2024). The structural approach to causal inference (Pearl, 2009; Pearl et al., 2016) provides a formal framework for defining counterfactuals. For instance, in the scenario described in Section 7.1.1, the approach begins by defining the *individual causal effect* (ICE) as the difference between each student’s potential outcomes, as in Equation 19.

$$\tau_i = Y_i \mid do(T_i = 1) - Y_i \mid do(T_i = 2) \quad (19)$$

where $do(T_i = t)$ represents the intervention operator, $Y_i \mid do(T_i = 1)$ represents the potential outcome under intervention $T_i = 1$, and $Y_i \mid do(T_i = 2)$ represents the potential outcome under intervention $T_i = 2$. Here, an *intervention* involves assigning a constant value to the treatment variable for each student’s potential outcomes. Note that if a student is assigned to intervention $T_i = 1$, the potential outcome under $T_i = 2$ becomes a counterfactual, as it is no longer observed nor observable. To address this challenge, the structural approach extends the ICE to the *average causal effect* (ACE, Equation 20), representing the average difference between the students’ observed potential outcomes and their counterfactual counterparts.

$$\begin{aligned}
\tau &= E[\tau_i] \\
&= E[Y_i \mid do(T_i = 1)] - E[Y_i \mid do(T_i = 2)]
\end{aligned} \tag{20}$$

Even though counterfactuals are unobservable, researchers can still identify the ACE from associational estimates by leveraging the structural approach. The approach identifies the ACE by statistically conditioning data on a *sufficient adjustment set* of variables X (Pearl, 2009; Pearl et al., 2016; Morgan and Winship, 2014). This *sufficient* set (potentially empty) must block all non-causal paths between T to Y without opening new ones. When such a set exists, then T and Y are *d-separated* by X ($T \perp Y \mid X$) (Pearl, 2009), and X satisfies the *backdoor criterion* (Neal, 2020, pp 37). Here, *conditioning* describes the process of restricting the focus to the subset of the population defined by the conditioning variable (Neal, 2020, pp. 32) (see Equation 21).

Conditioning on a sufficient adjustment set enables researchers to estimate the ACE, even when the data comes from an observational study. This process is feasible because such conditioning ensures that the ACE estimator satisfies several critical properties, including confounding elimination (Morgan and Winship, 2014). Naturally, the validity of claims about the causal effects of T on Y now hinges on the assumption that X serves as a sufficient adjustment set. However, as Kohler et al. (2019, pp. 150) noted, drawing conclusions about the real world from observed data inevitably requires assumptions. This requirement holds true for both observational and experimental data.

For instance, if researchers cannot conduct the randomized experiments described in Section 7.1.1 and must instead rely on observational data, they can still identify the ACE as long as an observed variable X , such as the socio-economic status of the school, satisfies the backdoor criterion. Under these circumstances, researchers first identify the *conditional average causal effect* (CACE, Equation 21)

$$CACE_t = E[Y_i \mid T_i = t, X] \tag{21}$$

From the CACE, researchers can identify the ACE from associational quantities as in Equation 22. This identification process is commonly known as the *backdoor adjustment*. Here, $E_X[\cdot]$ represents the marginal expected value over X (Morgan and Winship, 2014).

$$\begin{aligned}
\tau &= E[Y_i \mid do(T_i = 1)] - E[Y_i \mid do(T_i = 2)] \\
&= E_X[CACE_1 - CACE_2] \\
&= E_X[E[Y_i \mid T_i = 1, X] - E[Y_i \mid T_i = 2, X]]
\end{aligned} \tag{22}$$

Notably, the approach extends the ACE identification for a continuous variable T as in Equation 23, ensuring broad applicability across different causal scenarios (Neal, 2020, pp. 45)

$$\begin{aligned}
\tau &= E[Y_i \mid do(T_i = t)] \\
&= dE_X[E[Y_i \mid T_i = t, X]] / dt
\end{aligned} \tag{23}$$

7.1.3. Diving into the specifics

The structural approach to causal inference uses SCMs and DAGs to formally and graphically represent the presumed causal structure underlying the ACE (Pearl, 2009; Pearl et al., 2016; Gross et al., 2018; Neal, 2020). Essentially, these tools serve as *conceptual (theoretical) models* on which identification analysis rests (Schuessler and Selb, 2023, pp. 4). Thus, using these tools, researchers can determine which statistical models can identify (ACE, CACE, or other), assuming the depicted causal structure is correct (McElreath, 2020), thus enabling valid causal inference. Figure 9 shows the role of theoretical models in the inference process.

SCMs and DAGs support identification analysis through two key advantages. First, regardless of complexity, they can represent various causal structures using only five fundamental building blocks (Neal, 2020; McElreath, 2020). This feature allows researchers to decompose complex structures into manageable components, facilitating their analysis (McElreath, 2020). Second, they depict causal relationships in a non-parametric and fully interactive way. This flexibility enables feasible ACE identification strategies without defining the variables' data types, the functional form between them, or their parameters (Pearl et al., 2016, pp. 35).

Thus, Section 7.1.3.1 and Section 7.1.3.2 elaborate on the first advantage, while Section 7.1.3.2 and Section 7.1.3.3 do so for the second. Finally, Section 7.1.3.4 explains how researchers use SCMs and DAGs alongside Bayesian inference methods in the estimation process.

7.1.3.1. The five fundamental block for SCMs and DAGs.

Figures 10, 11, 12, 13, and 14 display the five fundamental building blocks for SCMs and DAGs. The left panels of the figures show the formal mathematical models, represented by the SCMs,

defined in terms of a set of *endogenous* variables $V = \{X_1, X_2, X_3\}$, a set of *exogenous* variables $E = \{e_{X1}, e_{X2}, e_{X3}\}$, and a set of functions $F = \{f_{X1}, f_{X2}, f_{X3}\}$ (Pearl, 2009; Cinelli et al., 2020). Endogenous variables are those whose causal mechanisms a researcher chooses to model (Neal, 2020). In contrast, exogenous variables represent *errors* or *disturbances* arising from omitted factors that the investigator chooses not to model explicitly (Pearl, 2009, pp. 27,68). Lastly, the functions, referred to as *structural equations*, express the endogenous variables as non-parametric functions of other variables. These functions use the symbol ‘:=’ to denote the asymmetrical causal dependence of the variables and the symbol ‘ \perp ’ to represent *d-separation*, a concept akin to (conditional) independence.

Notably, every SCM has an associated DAG (Pearl et al., 2016; Cinelli et al., 2020). The right panels of the figures display these DAGs. A DAG is a graph consisting of nodes connected by edges, where the nodes represent random variables. The term *directed* means that the edges extend from one node to another, with arrows indicating the direction of causal influence. The term *acyclic* implies that the causal influences do not form loops, ensuring the influences do not cycle back on themselves (McElreath, 2020). DAGs represent observed variables as solid black circles, while they use open circles for unobserved (latent) variables (Morgan and Winship, 2014). Although the *standard representation* of DAGs typically omits exogenous variables for simplicity, the *magnified representation* depicted in the figures offers one key advantage: including exogenous variables can help researchers highlight potential issues related to conditioning and confounding (Cinelli et al., 2020).



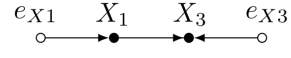
Figure 10: Two unconnected nodes

$$X_1 := f_{X1}(e_{X1})$$

$$X_3 := f_{X3}(X_1, e_{X3})$$

$$e_{X1} \perp e_{X3}$$

(a) SCM



(b) DAG

Figure 11: Two connected nodes or descendant

$$X_1 := f_{X1}(e_{X1})$$

$$X_2 := f_{X2}(X_1, e_{X2})$$

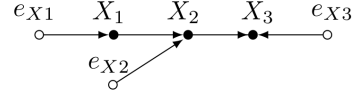
$$X_3 := f_{X3}(X_2, e_{X3})$$

$$e_{X1} \perp e_{X2}$$

$$e_{X1} \perp e_{X3}$$

$$e_{X2} \perp e_{X3}$$

(a) SCM



(b) DAG

Figure 12: Chain or mediator

$$X_1 := f_{X1}(X_2, e_{X1})$$

$$X_2 := f_{X2}(e_{X2})$$

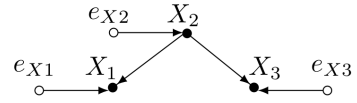
$$X_3 := f_{X3}(X_2, e_{X3})$$

$$e_{X1} \perp e_{X2}$$

$$e_{X1} \perp e_{X3}$$

$$e_{X2} \perp e_{X3}$$

(a) SCM



(b) DAG

Figure 13: Fork or confounder

$$X_1 := f_{X_1}(e_{X_1})$$

$$X_2 := f_{X_2}(X_1, X_3, e_{X_2})$$

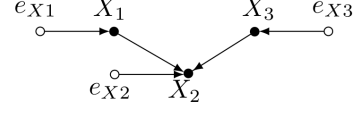
$$X_3 := f_{X_3}(e_{X_3})$$

$$e_{X_1} \perp e_{X_2}$$

$$e_{X_1} \perp e_{X_3}$$

$$e_{X_2} \perp e_{X_3}$$

(a) SCM



(b) DAG

Figure 14: Collider or immorality

A careful examination of these building blocks highlights the theoretical assumptions underlying their observed variables. SCM 10a and DAG 10b depict two unconnected nodes, representing a scenario where variables X_1 and X_3 are independent or not causally related. SCM 11a and DAG 11b illustrate two connected nodes, representing a scenario where a *parent* node X_1 exerts a causal influence on a *child* node X_3 . In this setup, X_3 is considered a *descendant* of X_1 . Additionally, X_1 and X_3 are described as *adjacent* because there is a *direct path* connecting them. SCM 12a and DAG 12b depict a *chain*, where X_1 influences X_2 , and X_2 influences X_3 . In this configuration, X_1 is a parent node of X_2 , which is a parent node of X_3 . This structure creates a *directed path* between X_1 and X_3 . Consequently, X_1 is an *ancestor* of X_3 , and X_2 fully *mediates* the relationship between the two. SCM 13a and DAG 13b illustrate a *fork*, where variables X_1 and X_3 are both influenced by X_2 . Here, X_2 is a parent node that *confounds* the relationship between X_1 and X_3 . Finally, SCM 14a and DAG 14b show a *collider*, where variables X_1 and X_3 are concurrent causes of X_2 . In this configuration, X_1 and X_3 are not causally related to each other but both influence X_2 (an *immorality*). Notably, all building blocks assume the errors are independent of each other and from all other variables in the graph, as evidenced by the pairwise relations $e_{X_1} \perp e_{X_2}$, $e_{X_1} \perp e_{X_3}$, and $e_{X_2} \perp e_{X_3}$.

Researchers can then use these building blocks to represent the scenario outlined in Section 7.1.2. SCM 15a and DAG 15b depict the plausible causal structure for this example. In this context, the variable X (socio-economic status of the school) is thought to be a confounder in the relationship between the teaching method T and the outcome Y . The figures display multiple descendant relationships such as $X \rightarrow T$, $X \rightarrow Y$, and $T \rightarrow Y$. They also highlight unconnected node pairs,

evident from the relationships $e_T \perp e_X$, $e_T \perp e_Y$, and $e_X \perp e_Y$. Additionally, the figures show one fork, $X \rightarrow \{T, Y\}$, and two colliders: $\{X, e_T\} \rightarrow T$ and $\{X, T, e_Y\} \rightarrow Y$.

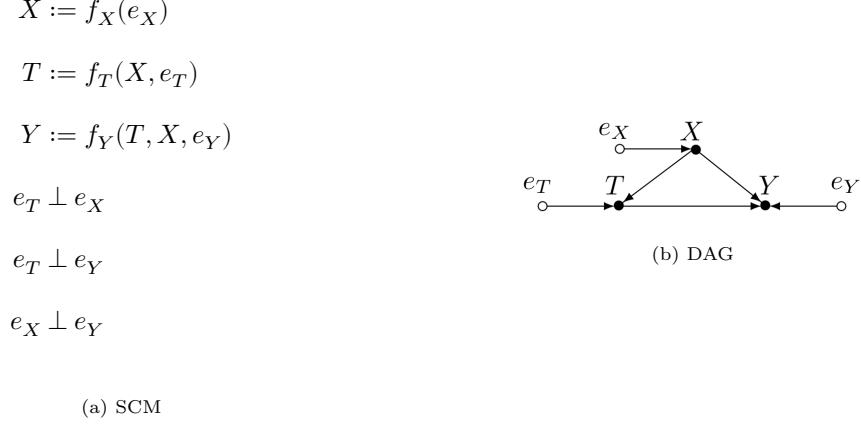


Figure 15: Plausible causal structure the scenario outlined in Section 7.1.2.

7.1.3.2. The probabilistic implications of these blocks.

Beyond their graphical capabilities, SCMs and DAGs can encode the probabilistic information embedded within a causal structure. They achieve this encoding by relying on three fundamental assumptions: the local Markov, the minimality, the causal edges assumption. The *local Markov assumption* encodes probabilistic independencies between variables by declaring that nodes in a graph are independent of all its non-descendants, given its parents (Neal, 2020, pp. 20). Meanwhile, the *minimality assumption* encodes probabilistic dependencies among variables by stating that every pair of adjacent nodes exhibits a dependency (Neal, 2020, pp. 21). Finally, the *causal edges assumption* encodes causal relationships between variables by declaring that each parent node acts as a direct cause of its children (Neal, 2020, pp. 22). Figure 16 illustrates how these assumptions influence the statistical and causal interpretations of graphs.

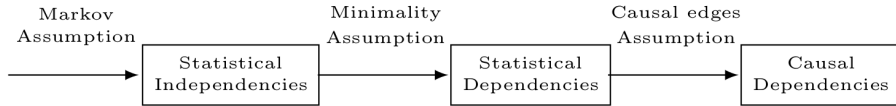


Figure 16: The flow of association and causation in graphs. Extracted and slightly modified from Neal (2020, pp. 31)

A notable implication of the assumptions underlying the probabilistic encoding is that any conceptual model described by an SCM and DAG can represent the joint distribution of variables more efficiently (Pearl et al., 2016, pp. 29). This expression takes the form of a product of conditional probability distributions (CPDs) of the type $P(\text{child} \mid \text{parents})$. This property is formally known

as the *Bayesian Network factorization* (BNF, Equation 24) (Pearl et al., 2016, pp. 29; Neal, 2020, pp. 21). In this expression, $pa(X_i)$ denotes the set of variables that are the parents of X_i .

$$\begin{aligned} P(X_1, X_2, \dots, X_P) &= X_1 \cdot \prod_{p=2}^P P(X_i | X_{i-1}, \dots, X_1) \quad (\text{by chain rule}) \\ &= X_1 \cdot \prod_{p=2}^P P(X_i | pa(X_i)) \quad (\text{by BNF}) \end{aligned} \tag{24}$$

This encoding enables researchers with conceptual (theoretical) knowledge in the form of an SCM and DAG to predict patterns of (in)dependencies in the data. As highlighted by Pearl et al. (2016, pp. 35), these predictions depend solely on the structure of these conceptual models without requiring the quantitative details of the equations or the distributions of the errors. Moreover, once researchers observe empirical data, the patterns of (in)dependencies in the data can provide significant insights into the validity of the proposed conceptual model.

The five fundamental building blocks described in Section 7.1.3.1 clearly illustrate which (in)dependencies can SMCs and DAGs predict. For instance, applying the BNF to the causal structure shown in the SCM 10a and DAG 10b enables researchers to express the joint probability distribution of the observed variables as $P(X_1, X_3) = P(X_1)P(X_3)$, supporting the theoretical assumption that the observed variables X_1 and X_3 are unconditionally independent ($X_1 \perp X_3$) (Neal, 2020, pp. 24). Conversely, when X_3 is unconditionally dependent on X_1 ($X_1 \not\perp X_3$), as depicted in the SCM 11a and DAG 11b, the BNF express their joint probability distribution as $P(X_1, X_3) = P(X_3 | X_1)P(X_1)$. Notably, these descriptions demonstrate the clear correspondence between the structural equations illustrated in Section 7.1.3.1 and the CPDs.

Beyond the insights gained from two-node structures, researchers can uncover more nuanced patterns of (in)dependencies from chains, forks, and colliders. These (in)dependencies apply to any data set generated by a causal model with those structures, regardless of the specific functions attached to the SCM (Pearl et al., 2016, pp. 36). For instance, applying the BNF to the chain structure depicted in the SCM 12a and DAG 12b allow researchers to represent the joint distribution for the observed variables as $P(X_1, X_2, X_3) = P(X_1)P(X_2 | X_1)P(X_3 | X_2)$. This expression implies that X_1 and X_3 are unconditionally dependent ($X_1 \not\perp X_3$), but conditionally independent when controlling for X_2 ($X_1 \perp X_3 | X_2$). Moreover, in the fork structure shown in the SCM 13a and DAG 13b, researchers can express the joint distribution of the observed variables as $P(X_1, X_2, X_3) = P(X_1 | X_2)P(X_2)P(X_3 | X_2)$. Similar to the chain structure, this expression allows researchers to further infer that X_1 and X_3 are unconditionally dependent ($X_1 \not\perp X_3$), but conditionally independent

dent when controlling for X_2 ($X_1 \perp X_3 \mid X_2$). Finally, researchers analyzing the collider structure illustrated in the SCM 14a and DAG 14b can express the joint distribution of the observed variables as $P(X_1, X_2, X_3) = P(X_1)P(X_2 \mid X_1, X_3)P(X_3)$. This representation allows researchers to infer that X_1 and X_3 are unconditionally independent ($X_1 \perp X_3$), but conditionally dependent when controlling for X_2 ($X_1 \not\perp X_3 \mid X_2$). The authors Pearl et al. (2016, pp. 37, 40, 41) and Neal (2020, pp. 25–26) provide the mathematical proofs for these conclusions.

Using these additional probabilistic insights, researchers can revisit the scenario in Section 7.1.2. In this context, applying the BNF to the SCM 17a structure, enables the representation of the joint probability distribution of the observed variables as $P(Y, T, X) = P(Y \mid T, X)P(T \mid X)P(X)$. From this expression, researchers can infer that the outcome Y is unconditionally dependent on the teaching method T ($Y \not\perp T$). This dependency arises from two key structures: a direct causal path from the teaching method T to the outcome Y , represented by the two-connected-nodes structure $T \rightarrow Y$ (black path in DAG 17b), and a confounding non-causal path from the teaching method T to the outcome Y through the socio-economic status of the school X , represented by the fork structure $T \leftarrow X \rightarrow Y$ (gray path in DAG 17b).

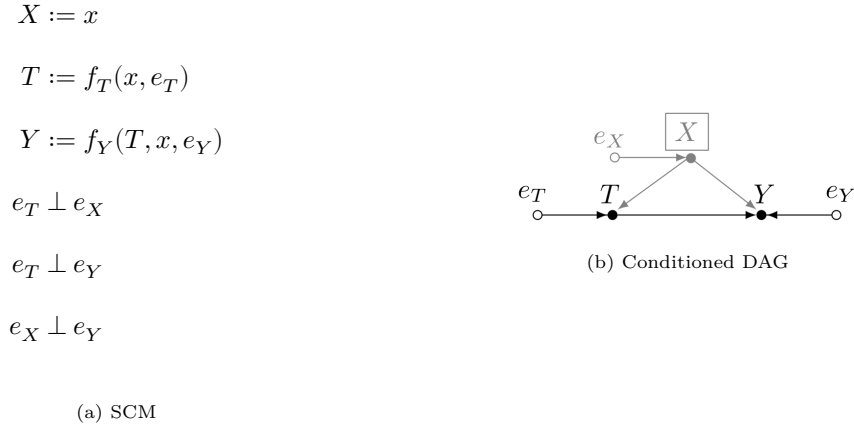


Figure 17: Plausible causal structure the scenario outlined in Section 7.1.2.

7.1.3.3. From probability to causality.

The structural approach to causal inference translates probabilistic insights into actionable strategies seeking to identify the ACE from associational quantities. The approach achieves this by relying on the *modularity assumption*, which posits that intervening on a node alters only the causal mechanism of that node, leaving others unchanged (Neal, 2020, pp. 34).

The modularity assumption underpins the concepts of manipulated graphs and Truncated Fac-

torization, which are essential for representing interventions $P(Y_i \mid do(T_i = t))$ within SCMs and DAGs. *Manipulated graphs* simulate physical interventions by removing specific edges from a DAG, while preserving the remaining structure unchanged (Neal, 2020, pp. 34). In parallel, *Truncated Factorization* (TF) achieves a similar simulation by removing specific functions from the conceptual model and replacing them with constants, while keeping the rest of the structure unchanged (Pearl, 2010). The probabilistic implications of this factorization are formalized in Equation 25, where S represents the subset of variables X_p directly influenced by the intervention, while an example illustrating these concepts follows below.

$$P(X_1, X_2, \dots, X_P \mid do(S)) = \begin{cases} \prod P(X_p \mid pa(X_p)) & \text{if } p \notin S \\ 1 & \text{otherwise} \end{cases} \quad (25)$$

Using the TF, researchers can define the *backdoor adjustment* to identify the ACE. This adjustment states that if a variable $X_p \in S$ serves as a *sufficient adjustment set* for the effect of X_a on X_b , then the ACE can be identified using Equation 26. The sufficient adjustment set (potentially empty) must block all non-causal paths between X_a and X_b without introducing new paths. If such a set exists, then X_a and X_b are *d-separated* by X_p ($X_a \perp X_b \mid X_p$) (Pearl, 2009), and X_p satisfies the *backdoor criterion* (Neal, 2020, pp. 37).

$$P(X_a \mid do(X_b = x)) = \sum_{X_p} P(X_a \mid X_b = x, X_p) P(X_p) \quad (26)$$

Ultimately, the backdoor adjustment enables researchers to express the ACE as:

$$\begin{aligned} \tau &= E[X_a \mid do(X_b = 1)] - E[X_a \mid do(X_b = 2)] \\ &= E_{X_p} [E[X_a \mid do(X_b = 1), X_p] - E[X_a \mid do(X_b = 2), X_p]] \\ &= \sum_{X_p} X_a \cdot P(X_a \mid X_b = 1, X_p) \cdot P(X_p) - \sum_{X_p} X_a \cdot P(X_a \mid X_b = 2, X_p) \cdot P(X_p) \end{aligned} \quad (27)$$

With these new insights, researchers revisiting the scenario in Section 7.1.3.2 can infer that the socio-economic status of the school, X , satisfies the backdoor criterion, assuming the causal structure depicted by the SCM 17a and DAG 17b is correct. This means that X serves as a sufficient adjustment set, as it effectively blocks all confounding non-causal paths introduced by the fork structure. Nevertheless, since Y remains dependent on T even after conditioning ($Y \not\perp T \mid X$), this

dependency can only be attributed to the direct causal effect $T \rightarrow Y$. Notably, for the purpose of identification, the conditioned DAG 17b is equivalent to the manipulated DAG 18b, because X satisfies the backdoor criterion.

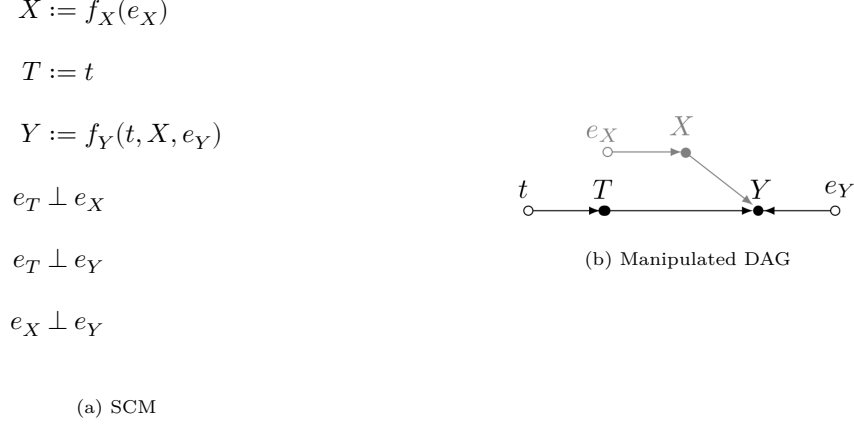


Figure 18: Plausible causal structure the scenario outlined in Section 7.1.3.2.

Researchers can then apply the *backdoor adjustment* to identify the ACE of T on Y . They achieve this by first identifying the CACE of T on Y by conditioning on X , and then marginalizing this effect over X to obtain the ACE. This process is expressed in Equation 28 (see Section 7.1.2).

$$\begin{aligned}
\tau &= E[Y_i \mid do(T_i = 1)] - E[Y_i \mid do(T_i = 2)] \\
&= E_X [E[Y_i \mid T_i = 1, X] - E[Y_i \mid T_i = 2, X]] \\
&= \sum_X Y_i \cdot P(Y_i \mid T_i = 1, X) \cdot P(X) - \sum_X Y_i \cdot P(Y_i \mid T_i = 2, X) \cdot P(X)
\end{aligned} \tag{28}$$

7.1.3.4. The estimation process.

Ultimately, researchers can use Bayesian inference methods to estimate the ACE. The approach begins by defining two probability distributions: the likelihood of the data, $P(X_1, X_2, \dots, X_P \mid \theta)$, and the prior distribution, $P(\theta)$ (Everitt and Skrongdal, 2010), where X_P represents a random variable, and θ represents a one-dimensional parameter space for simplicity. After observing empirical data, researchers can update the priors to posterior distributions using Bayes' rule in Equation 29:

$$P(\theta \mid X_1, X_2, \dots, X_P) = \frac{P(X_1, X_2, \dots, X_P \mid \theta) \cdot P(\theta)}{P(X_1, X_2, \dots, X_P)} \tag{29}$$

Given that the denominator on the right-hand side of Equation 29 serves as a normalizing constant

independent of the parameter θ , researchers can simplify the posterior updating process into three steps. First, they integrate new empirical data through the likelihood. Second, they update the parameters' priors to a posterior distribution according to Equation 30. Ultimately, they normalize these results to obtain a valid probability distribution.

$$P(\theta \mid X_1, X_2, \dots, X_P) \propto P(X_1, X_2, \dots, X_P \mid \theta) \cdot P(\theta) \quad (30)$$

Temporarily setting aside the definition of prior distributions $P(\theta)$, note that the posterior updating process depends heavily on the assumptions underlying the likelihood of the data. However, as the number of random variables, P , increases, this joint distribution quickly becomes intractable (Neal, 2020). This intractability is evident from Equation 31, where the likelihood distribution is expressed by multiple chained CPDs.

$$P(X_1, X_2, \dots, X_P \mid \theta) = P(X_1 \mid \theta) \prod_{p=2}^P P(X_p \mid X_{p-1}, \dots, X_1, \theta) \quad (31)$$

Nevertheless, researchers can manage the complexity of the likelihood by assuming specific local (in)dependencies among variables. SCMs and DAGs provide a formal framework to represent these assumptions, as detailed in Section 7.1.3.2. These assumptions improve model tractability and simplify the estimation process by enabling the derivation of the BNF of the likelihood (Equation 32). With this simplified structure, any probabilistic programming language can model the system and compute the parameter's posterior distribution using Equation 29.

$$P(X_1, X_2, \dots, X_P \mid \theta) = P(X_1 \mid \theta) \prod_{p=2}^P P(X_p \mid pa(X_p), \theta) \quad (32)$$

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