Evaluating Model Estimation Processes for Diagnostic Classification Models

By

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Evaluating Model Estimation Processes for Diagnostic Classification Models

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Diagnostic classification models (DCMs) are a class of models that define respondent ability on a set of predefined categorical latent variables. In recent years, the popularity of these models has begun to increase. As the community of researchers of practitioners of DCMs grow, it is important to examine the implementation of these models, including the process of model estimation. A key aspect of the estimation process that remains unexplored in the DCM literature is model reduction, or the removal of parameters from the model in order to create a simpler, more parsimonious model. The current study fills this gap in the literature by first applying several model reduction processes on a real data set, the Diagnosing Teachers’ Multiplicative Reasoning assessment (Bradshaw et al., 2014). Results from this analysis indicate that the selection of model reduction process can have large implications for the resulting parameter estimates and respondent classifications. A simulation study is then conducted to evaluate the relative performance of these various model reduction processes. The results of the simulation suggest that all model reduction processes are able to provide quality estimates of the item parameters and respondent masteries, if the model is able to converge. The findings also show that if the full model does not converge, then reducing the structural model provides the best opportunities for achieving a converged solution. Implications of this study and directions for future research are discussed.

Keywords. Diagnostic classification models, log-linear cognitive diagnosis model, model reduction, Monte Carlo simulation
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# Contents

1 Introduction 1

1.1 Study constraints .................................................. 2
1.2 Colophon ........................................................... 3

2 Literature Review 4

2.1 Structure of diagnostic classification models .................... 5
  2.1.1 The Q-matrix ................................................... 7
2.2 Types of DCMs ....................................................... 8
  2.2.1 Noncompensatory DCMs ....................................... 8
  2.2.2 Compensatory DCMs .......................................... 9
2.3 The log-linear cognitive diagnosis model ......................... 10
  2.3.1 LCDM measurement model .................................... 11
2.4 Structural models .................................................. 15
  2.4.1 Log-linear structural models ................................ 16
2.5 Model reduction .................................................... 17
  2.5.1 Model reduction in structural equation modeling .......... 18
  2.5.2 Model reduction in item response theory .................... 19
  2.5.3 Model reduction in DCMs ..................................... 20
2.6 The current study .................................................. 21

3 Pilot Study 23

3.1 Method ............................................................. 23
  3.1.1 DTMR data .................................................... 23
<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.1.2 Model estimation</td>
<td>25</td>
</tr>
<tr>
<td>3.2 Results</td>
<td>27</td>
</tr>
<tr>
<td>3.2.1 Measurement model results</td>
<td>27</td>
</tr>
<tr>
<td>3.2.2 Structural model results</td>
<td>32</td>
</tr>
<tr>
<td>3.3 Conclusions</td>
<td>34</td>
</tr>
<tr>
<td><strong>4 Methods</strong></td>
<td><strong>36</strong></td>
</tr>
<tr>
<td>4.1 Overview of Monte Carlo methods</td>
<td>36</td>
</tr>
<tr>
<td>4.2 The current simulation</td>
<td>37</td>
</tr>
<tr>
<td>4.2.1 Simulation conditions</td>
<td>37</td>
</tr>
<tr>
<td>4.2.2 Data generation process</td>
<td>38</td>
</tr>
<tr>
<td>4.2.3 Model reduction process</td>
<td>39</td>
</tr>
<tr>
<td>4.2.4 Outcome measures</td>
<td>41</td>
</tr>
<tr>
<td>4.2.5 Software</td>
<td>42</td>
</tr>
<tr>
<td><strong>5 Results</strong></td>
<td><strong>43</strong></td>
</tr>
<tr>
<td>5.1 Reduction by p-value</td>
<td>44</td>
</tr>
<tr>
<td>5.1.1 Convergence</td>
<td>44</td>
</tr>
<tr>
<td>5.1.2 Parameter recovery</td>
<td>45</td>
</tr>
<tr>
<td>5.1.3 Mastery classification</td>
<td>49</td>
</tr>
<tr>
<td>5.1.4 Model fit</td>
<td>53</td>
</tr>
<tr>
<td>5.1.5 Description of reduced parameters</td>
<td>56</td>
</tr>
<tr>
<td>5.2 Reduction by heuristic</td>
<td>61</td>
</tr>
<tr>
<td>5.2.1 Convergence</td>
<td>62</td>
</tr>
<tr>
<td>5.2.2 Parameter recovery</td>
<td>64</td>
</tr>
<tr>
<td>5.2.3 Mastery classification</td>
<td>67</td>
</tr>
<tr>
<td>5.2.4 Model fit</td>
<td>71</td>
</tr>
<tr>
<td>5.2.5 Description of reduced parameters</td>
<td>74</td>
</tr>
</tbody>
</table>
6 Conclusions

6.1 Limitations and future directions ................................. 80

References 82

A Parameter Recovery with Reduction from P-values 91

A.1 Individual Bias ..................................................... 91
A.2 Individual Mean Square Error ................................. 97

B Parameter Recovery with Reduction from Heuristic 103

B.1 Individual Bias ..................................................... 103
B.2 Individual Mean Square Error ................................. 109
## List of Figures

3.1 Flowchart of model reduction processes ........................................ 26
3.2 DTMR respondent classification under each model reduction process ........ 34
4.1 Flowchart of simulation study reduction processes .............................. 40
5.1 Convergence rates when reducing using p-values ............................... 45
5.2 Bias in measurement model main effect estimates when reducing using p-values . 46
5.3 Mean square error in measurement model main effect estimates when reducing using p-values ................................................................. 47
5.4 Bias in structural model parameter estimates when reducing using p-values . 48
5.5 Mean square error in structural model parameter estimates when reducing using p-values ................................................................. 49
5.6 Average correct classification rate of attribute mastery when reducing using p-values 50
5.7 Average Cohen’s $\kappa$ of attribute mastery when reducing using p-values ... 51
5.8 Average correct classification rate of profile assignment when reducing using p-values ................................................................. 52
5.9 Average Cohen’s $\kappa$ of profile assignment when reducing using p-values ... 53
5.10 Number of selections as best fitting model as measured by the AIC when reducing using p-values ................................................................. 54
5.11 Number of selections as best fitting model as measured by the BIC when reducing using p-values ................................................................. 55
5.12 Number of selections as best fitting model as measured by the adjusted BIC when reducing using p-values ................................................................. 56
5.13 Proportion of correct measurement model reductions when reducing with p-values . 57
5.14 Proportion of correct structural model reductions when reducing with p-values . . . 58
5.15 Distributions of estimates and standard errors for reduced measurement model pa-
arameters when reducing with p-values . . . . . . . . . . . . . . . . . . . . . . . . . 60
5.16 Distributions of estimates and standard errors for reduced structural model param-
eters when reducing with p-values . . . . . . . . . . . . . . . . . . . . . . . . . . 61
5.17 Convergence rates when reducing using a heuristic . . . . . . . . . . . . . . . . 63
5.18 Bias in measurement model main effect estimates when reducing using a heuristic . 64
5.19 Mean square error in measurement model main effect estimates when reducing
using a heuristic . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 65
5.20 Bias in structural model parameter estimates when reducing using a heuristic . . . 66
5.21 Mean square error in structural model parameter estimates when reducing using a
heuristic . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 67
5.22 Average correct classification rate of attribute mastery when reducing using a heuris-
tic . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 68
5.23 Average Cohen’s κ of attribute mastery when reducing using a heuristic . . . . . 69
5.24 Average correct classification rate of profile assignment when reducing using a
heuristic . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 70
5.25 Average Cohen’s κ of profile assignment when reducing using a heuristic . . . . . 71
5.26 Number of selections as best fitting model as measured by the AIC when reducing
using a heuristic . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 72
5.27 Number of selections as best fitting model as measured by the BIC when reducing
using a heuristic . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 73
5.28 Number of selections as best fitting model as measured by the adjusted BIC when
reducing using a heuristic . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . . 74
5.29 Proportion of correct measurement model reductions when reducing with a heuristic 75
5.30 Proportion of correct structural model reductions when reducing with a heuristic . . 76
A.1 Bias in measurement model intercept estimates when reducing using p-values . . . 91
A.2 Bias in measurement model main effect estimates when reducing using p-values . . 92
A.3 Bias in measurement model 2-way interaction estimates when reducing using p-
values ................................................................. 93
A.4 Bias in measurement model 3-way interaction estimates when reducing using p-
values ................................................................. 94
A.5 Bias in measurement model 4-way interaction estimates when reducing using p-
values ................................................................. 95
A.6 Bias in structural model estimates when reducing using p-values ................. 96
A.7 MSE in measurement model intercept estimates when reducing using p-values . 97
A.8 MSE in measurement model main effect estimates when reducing using p-
values ................................................................. 98
A.9 MSE in measurement model 2-way interaction estimates when reducing using p-
values ................................................................. 99
A.10 MSE in measurement model 3-way interaction estimates when reducing using p-
values ................................................................. 100
A.11 MSE in measurement model 4-way interaction estimates when reducing using p-
values ................................................................. 101
A.12 MSE in structural model estimates when reducing using p-values ............... 102
B.1 Bias in measurement model intercept estimates when reducing using a heuristic . 103
B.2 Bias in measurement model main effect estimates when reducing using a heuristic . 104
B.3 Bias in measurement model 2-way interaction estimates when reducing using a heuristic ................................................................. 105
B.4 Bias in measurement model 3-way interaction estimates when reducing using a heuristic ................................................................. 106
B.5 Bias in measurement model 4-way interaction estimates when reducing using a heuristic ................................................................. 107
B.6 Bias in structural model estimates when reducing using a heuristic ............. 108
B.7 MSE in measurement model intercept estimates when reducing using a heuristic . 109
B.8 MSE in measurement model main effect estimates when reducing using a heuristic 110
B.9 MSE in measurement model 2-way interaction estimates when reducing using a
        heuristic ................................................................. 111
B.10 MSE in measurement model 3-way interaction estimates when reducing using a
        heuristic ................................................................. 112
B.11 MSE in measurement model 4-way interaction estimates when reducing using a
        heuristic ................................................................. 113
B.12 MSE in structural model estimates when reducing using a heuristic ............. 114
List of Tables

2.1 Example cross classification ................................................. 10
2.2 Example cross classification with latent trait .......................... 11
2.3 Log-linear structural model for a 2-attribute assessment .......... 17

3.1 DTMR Q-matrix ................................................................. 24
3.2 DTMR estimates of item intercepts, $\lambda_{i,0}$ ........................ 28
3.3 DTMR estimates of item main effects for Referent Units, $\lambda_{i,1,(1)}$ 29
3.4 DTMR estimates of item main effects for Partitioning and Iterating, $\lambda_{i,1,(2)}$ 29
3.5 DTMR estimates of item main effects for Appropriateness, $\lambda_{i,1,(3)}$ 30
3.6 DTMR estimates of item main effects for Multiplicative Comparison, $\lambda_{i,1,(4)}$ 30
3.7 DTMR estimates of item interactions between Referent Units and Partitioning and Iterating, $\lambda_{i,2,(1,2)}$ 31
3.8 DTMR estimates of item interactions between Referent Units and Multiplicative Comparison, $\lambda_{i,2,(1,4)}$ 31
3.9 DTMR estimates of item interactions between Partitioning and Iterating and Multiplicative Comparison, $\lambda_{i,2,(2,4)}$ 31
3.10 DTMR estimates of item interactions between Appropriateness and Multiplicative Comparison, $\lambda_{i,2,(3,4)}$ 32
3.11 DTMR estimates of structural parameters ............................ 33

5.1 Saturated model convergence rates ................................. 43
Chapter 1

Introduction

Over the past several years, diagnostic classification models (DCMs) have become a more prominent research focus in the field of educational assessment, and psychometrics more broadly (Bradshaw, 2017; Rupp & Templin, 2008b; Rupp, Templin, & Henson, 2010). Rather than providing a single scaled-score for unidimensional construct, as is common in many item response theory based assessments (see Ayala, 2009), DCMs are multidimensional assessments that provide as their scores a profile of mastery or non-mastery on the skills, or attributes, that are assessed. Thus, DCMs are able to provide more detailed and actionable information about the skills a student has mastered, and the skills that could use more instruction.

In all multidimensional models (e.g., multidimensional item response theory, structural equation modeling, DCMs), there is a measurement model that relates observed data to the latent traits and a structural model that defines the relationships between the latent traits. If all possible parameters are estimated for both the measurement and structural models, then it is likely that unnecessary parameters are estimated, which can impact the stability of other parameters and scores, as well as increasing the complexity and intensity of computation (Browne, Rockloff, & Rawat, 2016). However, Templin & Bradshaw (2014b) note that it is also important to estimate enough parameters to capture the full complexity of the data. Model reduction refers to the process of removing parameters to provide a more parsimonious model while still capturing the appropriate level of complexity.

In the context of DCMs, there is little research or guidance as to how the model reduction process should take place. For instance, the measurement model and structural model could be reduced simultaneously, one could be reduced after the other, or only one could be reduced. An
exploration of these various processes using an assessment known as Diagnosing Teachers’ Multiplicative Reasoning assessment (Chapter 3) shows that the choice of model reduction process can have a profound impact on the final set of parameters included in the model, the estimates and standard errors of the parameters across processes, and respondent assignment to attribute profiles.

The current study further explores this gap in the literature concerning best practices for model reduction of DCMs. A simulation study is conducted, whereby data is simulated from the log-linear cognitive diagnosis model (section 2.3), and then the DCM is estimated using each of the possible model reduction processes. Bias and mean-squared error of the parameter estimates, along with estimated attribute mastery agreement provide insight as to which model reduction process is most appropriate under a variety of data generation conditions. The findings of this study have practical implications for the estimation of DCMs, as the simulation study provides evidence for effectiveness of various model reduction processes. Additionally, practitioners using DCMs in an applied setting will be able to benefit, as a more parsimonious model that is still accurate may provide a more efficient estimation process.

1.1 Study constraints

Although DCMs can be estimated with attributes that have more than one latent category (Rupp et al., 2010), this paper limits the discussion to binary latent attributes. Binary attributes are the most commonly used with DCMs, and this limitation simplifies the problem for the initial investigation proposed in this study. Further, the proposed study limits the discussion of data to dichotomously scored items. There are generalized DCMs that can accommodate alternative response types (e.g., Templin, Henson, Rupp, Jang, & Ahmed, 2008), however, these have not been widely used in the literature. Thus, the proposed study is limited to the types of DCMs that have been most widely investigated and used operationally: binary attributes with dichotomous item responses.
1.2 Colophon

This document was written in Rmarkdown inside RStudio (RStudio, 2018) using the *rmarkdown* (Allaire et al., 2017), *bookdown* (Xie, 2017a), and *jayhawkdown* (Thompson & Johnson, 2017) packages. The raw Rmarkdown was converted to html and pdf documents using pandoc (“Pandoc,” 2017) and the *knitr* package (Xie, 2017b). All graphics were created using the *ggplot2* (Wickham & Chang, 2018), *ggforce* (Pedersen, 2016), and *colorblindr* (McWhite & Wilke, 2018) packages, and tables were formatted using the *kableExtra* package (Zhu, 2018). The website was made with jekyll (Preston-Werner, 2018) and published to Netlify (Netlify, 2018) with Travis-CI (Travis CI, 2018). The source code for this document is available on GitHub.
Chapter 2

Literature Review

Diagnostic classification models (DCMs; also known as cognitive diagnostic models), are class of psychometric models that define a mastery profile on a predefined set of attributes (Rupp & Templin, 2008b; Rupp et al., 2010). These attributes are categorical in nature, and although they can consist of more than two categories, they most usually are binary (Bradshaw, 2017). Given an attribute profile for an individual, the probability of providing a correct response to an item is determined by the attributes that are required by the item.

This profile of attribute mastery that gives rise to item responses makes DCMs an inherently different type of assessment than what is most commonly used in psychometrics. For example classical test theory (DeVellis, 2006), item response theory (Reckase, 2009), and structural equation modeling (Ullman & Bentler, 2003) all assume a continuous latent trait. This can result in difficulty in interpreting what an assessment score means. In an educational setting, a process known as standard setting (Cizek, 2006) is typically conducted to categorize the continuous score so that stakeholders and parents can better interpret what a score means (Hambleton, 2006). In contrast, assessments that are scaled with a diagnostic model provide a categorical class for each attribute that is mastered. This allows for a greater differentiation of respondent latent traits. However, a decision must still be made as to what probability of attribute mastery is sufficient for reporting an individual as a master.

Take, for example, a standard K-12 math assessment. Using traditional test scaling methods, a student would receive an overall math score, performance level determined by the standard setting process, and possibly a selection of subscores. However, because the unidimensional variants of these methods are most commonly used, subscores have been shown to be problematic with these
types of assessments (Feinberg & Wainer, 2014; Sinharay, Haberman, & Wainer, 2011).

Diagnostic models on the other hand are multidimensional models. Thus, if the math assessment were scaled using diagnostic models, the student would receive a probability of mastery on each of the attributes that was assessed (Bradshaw & Templin, 2014), although it is possible to also use a standard setting process within DCMs if desired (see Templin, 2010; Templin, Poggio, Irwin, & Henson, 2007). What these attributes are is determined in the test design process. They could be specific skills, educational standards, or subareas within the larger content area (e.g., algebra, geometry, and statistic all fall within the larger math construct). Therefore, it is critical to determine what level of score reporting is desired prior to test construction. For example both Rupp et al. (2010) and Almond, Mislevy, Steinberg, Yan, & Williamson (2015) discuss the evidence centered design framework, and how this approach to validity can aid in the construction of a diagnostic assessment. Under the evidence centered design framework, generally speaking, test design begins with the inferences about student ability that are desired, and then works back to the evidence needed to support those inferences. In this way, the grain size of the desired inferences will dictate the grain size of the attribute definitions.

In this chapter, the statistical structure of DCMs is outlined, the key differences between traditional sub-types of DCMs are highlighted. The log-linear cognitive diagnosis model is then explored in more depth, as this model subsumes all other DCMs and is the basis for this study. Finally, model estimation and reduction techniques are discussed across a variety of psychometric models, including diagnostic models, item response theory, and structural equation modeling.

2.1 Structure of diagnostic classification models

In this paper, the discussion of diagnostic models is restricted to binary attributes assessed by dichotomously scored items. Practically, this means that all models presented are extensions of latent class models. Specifically, DCMs can be thought of as a restricted latent class model where each class represents a profile of attribute mastery. When using binary attributes, the number of unique classes is equal to $2^A$, where $A$ is the number of attributes assessed. Given the specification
of available attribute profiles, the probability of respondent \( r \) providing a response to an item is as follows.

\[
P(X_r = x_r) = \sum_{c=1}^{C} \nu_c \prod_{i=1}^{I} \pi_{ic}^{x_{ir}} (1 - \pi_{ic})^{1-x_{ir}}
\]

(2.1)

In equation (2.1) \( \pi_{ic} \) is the probability of a respondent in class \( c \) providing a correct response to item \( i \), and \( x_{ir} \) is the response (i.e., 0, 1) of respondent \( r \) to item \( i \). Thus, \( \pi_{ic}^{x_{ir}} (1 - \pi_{ic})^{1-x_{ir}} \) can be described in words as the probability of a respondent in class \( c \) providing the observed response to item \( i \). The probabilities are then multiplied across all items, giving, the probability of a respondent in class \( c \) providing the observed response pattern. This portion of equation (2.1) is known as the \textit{measurement model}, and defines how the items are related to the attributes (equation (2.2) shows just the measurement model).

\[
\prod_{i=1}^{I} \pi_{ic}^{x_{ir}} (1 - \pi_{ic})^{1-x_{ir}}
\]

(2.2)

Continuing with equation (2.1), the probability of respondent in class \( c \) providing the observed response vector is then by multiplied by \( \nu_c \), the probability that any given respondent belongs to class \( c \). This product represents the probability that a given respondent is in class \( c \) and provided the observed response pattern. Summing over all possible classes gives the probability that a randomly chosen respondent would provide the observed response pattern. This section of equation (2.1) that defines the joint probability of membership in each class is known as the \textit{structural model} (equation (2.3) shows just the structural model). In the structural \( \nu \) is constrained to sum to 1, such that the probability of a respondent not belonging to any class is 0.

\[
\sum_{c=1}^{C} \nu_c
\]

(2.3)

Historically, diagnostic models have used an unconstrained structural model, meaning that the values \( \nu \) directly correspond to the proportion of respondents estimated to be in each of the latent classes. Thus, the structural model is consistent across the wide variety of diagnostic models that
exists. What differs between these models is how the measurement model, or how the items relate to the attributes. This process begins with the specification of a Q-matrix.

### 2.1.1 The Q-matrix

The specification of which attributes are measured by each item is defined *a priori* by the Q-matrix. The Q-matrix is an $n \times a$ matrix filled with 0s and 1s. A 0 indicates the item is not measured by the attribute, whereas a 1 indicates that the item is measured by the attribute (Tatsuoka, 1983). The Q-matrix is developed in consultation with content area experts to determine the attributes that need to be present in order for the item to be answered correctly (Bradshaw, 2017). Because the Q-matrix defines how the items relate to the latent attributes, the correct specification of the Q-matrix is critical to the accuracy of the parameter estimates and scores. Both Kunina-Habenicht, Rupp, & Wilhelm (2012) and Rupp & Templin (2008a) used simulation studies to demonstrate the ill-effects of misspecification on classification accuracy and parameter bias. Given this importance, it is common practice to make changes to the Q-matrix following the estimation of the DCM (Rupp et al., 2010).

For example de la Torre (2008) proposed a method for empirically validating the Q-matrix following estimation with the deterministic-input, noisy-and-gate model (see section 2.2.1). Using this method, de la Torre found acceptable Type I and Type II error rates, indicating that the method was able to adequately identify places where the Q-matrix was both correctly and incorrectly specified. Similarly, DeCarlo (2011) found that changing the Q-matrix structure could significantly improve the placement of respondents into latent classes. For example, the initial specification of the Q-matrix for fraction subtraction data (Tatsuoka, 1990) leads to respondents with no correct answers mastering the majority of skills. By changing the specification of the Q-matrix, DeCarlo (2011) was able to correct this, resulting in a more interpretable output. Chen, Liu, Xu, & Ying (2015) took this approach to the extreme by estimating the entire Q-matrix based only on the dependencies seen in item responses. Using this method, content experts are removed from the process of creating the Q-matrix entirely, and it is specified entirely by empirical methods.
What the Q-matrix is unable to define is how the attributes interact with each other on a given item to influence performance. If an item is measured by multiple attributes, does the respondent have to have mastered all of the attributes in order to have a high probability of answering the item correctly? Or would mastery of any of the attributes be sufficient? This definition of how the attributes interact with the items is defined by the measurement model (equation (2.2)). Traditionally, this choice of compensatory versus non-compensatory has been accomplished by choosing one of a variety of DCMs that have been proposed in the literature.

2.2 Types of DCMs

Traditionally, the type of compensation employed in the measurement model has been defined through the selection of a specific DCM. The individual types of DCMs each defined the compensatory or non-compensatory nature of the attributes and items differently. Thus, the relationships of attributes and items must be assumed *a priori*, and then enforced by the selected model. This relationship can be either non-compensatory or compensatory. Ostensibly, both compensatory and non-compensatory models could be estimated, compared, and then a final model selected *a posteriori*; however, usually when selecting one of these models, there is a conceptual reason for the selection, which may not be compatible with other types of DCMs. Non-compensatory DCMs require all of the attributes measured by an item to be mastered in order for the item to be answered correctly. In compensatory DCMs, mastery of some of the attributes measured by an item may be enough to provide a high probability of success. A high level description of these classes of DCMs follows.

2.2.1 Noncompensatory DCMs

Non-compensatory DCMs are defined such that all attributes associated with an item must be mastered in order for the respondent to have a high probability of answering the item correctly. In other words, having an excess of ability on one of the attributes measured by an item cannot make up
for the lack of ability on another. This class of DCMs includes the determinisitic-input, noisy-and-gate (DINA; de la Torre & Douglas, 2004; Haertel, 1989; Junker & Sijtsma, 2001), noisy-input, deterministic-and gate (NIDA; Henson & Douglas, 2005; Junker & Sijtsma, 2001), and reduced non-compensatory reparameterized unified (reduced NC-RUM; DiBello, Stout, & Roussos, 1995; Hartz, 2002) models. In the DINA and NIDA models, there are slipping and guessing parameters that are held constant across items or attributes respectively. In these models, the slipping parameter represents the probability of incorrectly applying an attribute that has be mastered, whereas the guessing parameter represents the probability of correctly applying an attribute that hasn’t been mastered. The reduced NC-RUM is parameterized slightly differently. In this model, the probability of providing a correct response when all required attributes have been mastered, with a penalty factor then applied for each attribute that isn’t mastered. However, in all of these models, the presence of one of the required attributes is unable to make up for the absence of another.

### 2.2.2 Compensatory DCMs

In contrast to the non-compensatory DCMs outlined above, compensatory DCMs are structured such that mastering a subset of the required attributes is sufficient to provide a correct response to the item. This means that not only a subset attributes that are measured by an item have to be mastered in order for the respondent to have a high probability of success. DCMs in this class include the deterministic-input, noisy-or-gate (DINO; Templin & Henson, 2006), noisy-input, deterministic-or-gate (NIDO; Rupp & Templin, 2008b), compensatory reparameterized unified (C-RUM; Hartz, 2002) models. The DINO model is parameterized similarly to the DINA and NIDA models, with slipping and guessing parameters that are held constants across items. However, in this model, the slipping parameter represents the probability of providing an incorrect response when **at least one** of the required attributes has been mastered (rather than when all attributes have been mastered as in the DINA model). A similar interpretation is made for the guessing parameter.

The NIDO and C-RUM models are parameterized slightly differently. Rather than modeling parameters on the probability scale, a linear predictor is estimated on the log-odds scale, and then
mapped to item scores using the logit link function (see section 2.3.1). In the NIDO model, an intercept is added to the linear predictor for all attributes measured by the item, and an additional main effect parameter for each of the mastered attributes. The C-RUM model is similar; however, rather than an estimating intercept for each attribute, the C-RUM model estimates an intercept for the entire item, which represents the log-odds of a correct response when none of the measured attributes are mastered. An additional main effect term is then added for each of the mastered attributes.

### 2.3 The log-linear cognitive diagnosis model

The log-linear cognitive diagnosis model (LCDM) is a general framework for diagnostic models that subsumes most of the existing DCMs, including those discussed in section 2.2 (Henson, Templin, & Willse, 2008; Rupp et al., 2010). Log-linear models are most commonly used in categorical data analysis when examining the change in frequency of respondents in a category across groups (Agresti, 2012). In these models, the frequency of respondents in a category is predicted by dummy coded grouping variables. Consider an example where a researcher is attempting to determine if there is a relationship between gender and political party affiliation (for the purposes of this example this will be limited to democratic or republican). This would result a 2x2 table similar to Table 2.1.

<table>
<thead>
<tr>
<th></th>
<th>Democratic</th>
<th>Republican</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>400</td>
<td>500</td>
</tr>
<tr>
<td>Female</td>
<td>600</td>
<td>300</td>
</tr>
</tbody>
</table>

The relationship between gender and party affiliation would be written mathematically as:

$$\ln(F_{ij}) = \lambda_0 + \lambda_i^{Gender} + \lambda_j^{Party} + \lambda_{ij}^{Gender\times Party}$$ (2.4)

In equation (2.4) the log frequency of respondents in a cell is given by a linear predictor. The
intercept, $\lambda_0$ represents the frequency for individuals in the reference group for both gender and party affiliation. The next two terms, $\lambda_i^{\text{Gender}}$ and $\lambda_j^{\text{Party}}$ represent the simple main effects for gender and party affiliation respectively. Finally, the interaction term, $\lambda_{ij}^{\text{Gender} \times \text{Party}}$ represents how related the two factors are. For instance, if gender and party affiliation are completely independent of one another, the interaction term would be equal to 0.

The LCDM is a log-linear model with categorical latent traits. Consider an item on an achievement test that measures a single attribute, $\alpha_1$. This would lead to a cross classification table similar to Table 2.1, but with a latent attribute.

Table 2.2: Example cross classification with latent trait

<table>
<thead>
<tr>
<th></th>
<th>Master</th>
<th>Nonmaster</th>
</tr>
</thead>
<tbody>
<tr>
<td>Correct (X=1)</td>
<td>900</td>
<td>200</td>
</tr>
<tr>
<td>Incorrect (X=0)</td>
<td>100</td>
<td>600</td>
</tr>
</tbody>
</table>

The mathematical definition of Table 2.2 would be as follows:

$$\ln(F_{ij}) = \lambda_0 + \lambda_i^{\alpha_1} + \lambda_j^x + \lambda_{ij}^{\alpha_1 \times x} \quad (2.5)$$

Table 2.2 and equation (2.5) could both be extended to multiple latent attributes by creating a three way cross classification table and adding the appropriate main effects and additional interaction terms. Because mastery of the attributes is unobserved, it must be estimated by relating the observed response to the unobserved attribute. As discussed in section 2.2, the relationship between observed data and the latent attributes is known as the measurement model.

### 2.3.1 LCDM measurement model

In order to use a log-linear model to predict probabilities of events occurring (rather than frequencies), a different link function must be used. Equations (2.4) and (2.5) used a log-link, as frequencies are only bounded on the lower end of the distribution by 0. Probabilities, on the other hand, are bounded by 0 on the lower and 1 on the upper ends of the distribution. Therefore, a lo-
gistic, or logit, link is used. As discussed in section 2.2.2, these types of generalized linear models involve combining the predictors into what is known as a kernel (or linear predictor in the generalized linear modeling literature; Stroup, 2012), which is an unbounded continuous value that is mapped to the item responses through a link function. When dealing with dichotomous data, this is most commonly achieved using the logit link function (Stroup, 2012), defined in equation (2.6).

\[ \eta_{ic} = \text{logit}(\pi_{ic}) = \ln \left( \frac{\pi_{ic}}{1 - \pi_{ic}} \right) \]  

(2.6)

Similarly, the inverse of the logit can be expressed as follows.

\[ \pi_{ic} = \text{logit}^{-1}(\eta_{ic}) = \frac{\exp(\eta_{ic})}{1 + \exp(\eta_{ic})} \]  

(2.7)

The inverse logit in equation (2.7) is more commonly seen in psychometrics, especially in reference to item response theory (Ayala, 2009).

For the LCDM, the general notation used by Rupp et al. (2010) for parameters in the linear predictor is \( \lambda_{i,1,(a,a',...)} \). In this notation, the first subscript identifies the item for the parameter, the second parameter indicates the level of the parameter (i.e., 0 = intercept, 1 = main effect, 2 = two-way interaction, etc.), and the third subscript specifies which attribute(s) are measured by the parameter. For example, if item 1 on an assessment measured both attributes 1 and 2, the probability of a correct response would be defined by an intercept, \( \lambda_{0} \), a simple main effect for attribute 1, \( \lambda_{1,1,(1)} \), a simple main effect for attribute 2, \( \lambda_{1,1,(2)} \), and the interaction between attributes 1 and 2 \( \lambda_{1,2,(1,2)} \).

For any number of attributes, A, the kernel for the logit can be defined as follows:

\[ \text{kernel}_{i} = \lambda_{i,0} + \sum_{a=1}^{A} \lambda_{i,1,(a)} \alpha_{ca}q_{ia} + \sum_{a=1}^{A} \sum_{a'=1}^{A} \lambda_{i,2,(a,a')} \alpha_{ca} \alpha_{ca'}q_{ia}q_{ia'} + \ldots \]  

(2.8)

Equation (2.8) demonstrates that the kernel for item is made up of the intercept, all main effects for attributes that have both been mastered by individuals in latent class \( c \) and are measured by item \( i \), and all two-way interactions that meet the conditions of all attributes in the interaction have been
mastered by latent class $c$ are measured by item $i$. Equation (2.8) could continue on, adding higher level interaction terms as more and more attributes are measured by item $i$, up to the total number of attributes, $A$.

Written more succinctly, equation (2.8) can be expressed with matrix notation as:

$$\text{kernel}_i = \lambda_{i,0} + \lambda_i^T h(\alpha_c, q_i)$$

(2.9)

For item $i$, $\lambda_i^T$ represents the transpose of the $(2^A - 1) \times 1$ vector of item parameters that contains the main effects and interactions, and $h(\alpha_c, q_i)$ is the $(2^A - 1) \times 1$ vector of attribute and Q-matrix combinations. Thus, written in a more general form, the probability of a respondent in latent class $c$ providing a correct response to item $i$ can be defined as:

$$\pi_{ic} = P(X_{ic} = 1 | \alpha_c) = \frac{\exp(\lambda_{i,0} + \lambda_i^T h(\alpha_c, q_i))}{1 + \exp(\lambda_{i,0} + \lambda_i^T h(\alpha_c, q_i))}$$

(2.10)

This expression of a DCM has many advantages over those defined in section 2.2. First, the LCDM has parameters that are easier to interpret than those seen in the traditional DCMs. For example, the DINA and NIDA models both contain guessing and slipping parameters, but the interpretation of them differs due to how parameters are constrained in these models. Additionally, the slipping parameter (the probability of getting the item wrong despite having mastered all of the constituent attributes) is less useful than $(1 - s_i)$, or the probability of providing a correct response, which is usually the value of interest. In contrast, the LCDM parameters are directly analogous to the parameters of a generalized linear models. Each parameter represents the change in the log-odds of a correct response.

Additionally, by placing constraints on the parameters of the LCDM, it is possible to estimate the aforementioned DCMs. For example, the DINA model requires that all attributes be mastered in order to increase the probability of a correct response. This can be accomplished by constraining all parameters except the intercept and highest level interaction term of equation (2.8) to be 0 (Henson et al., 2008; Rupp et al., 2010). With these constraints, the log-odds of success are equal
to $\lambda_{i,0}$ when not all attributes have been mastered, and $\lambda_{i,0} + \lambda_{i,2,(a,a')}$ when all attributes have been mastered (assuming the item only measures two attributes). The inverse logit (equation (2.7)) of $\lambda_{i,0}$ is equal to the guessing parameter $g_i$ in the DINA model, and the inverse logit of $\lambda_{i,0} + \lambda_{i,2,(a,a')}$ is equal to $(1 - s_i)$.

Similarly, the DINO model can also be replicated through constraints on the LCDM model. In the DINO model, mastering one attribute is just as good as mastering a different or multiple attributes that are measured by the item. Thus, the first constraint is that the main effects must be equal. If an item measures two attributes, the increase in the log-odds of providing a correct response is equal, regardless of which of the two is mastered. The second constraint is that the interaction term is equal to the negative of the main effect parameter. This means that three parameters (two main effects and an interaction) all have the same absolute value, but the main effects are positive and the interaction is negative. This has the effect of the interaction cancelling out the additional increase in log-odds of a correct response for mastering additional attributes. Thus, the kernel for LCDM parameterization of the DINO model can be written as:

$$\text{kernel}_i = \lambda_{i,0} + \lambda_i \alpha_1 + \lambda_i \alpha_2 - \lambda_i \alpha_1 \alpha_2$$

$$= \lambda_{i,0} + \lambda_i (\alpha_1 + \alpha_2 - \alpha_1 \alpha_2) \tag{2.11}$$

When neither of the attributes measured by the item have been mastered, $\alpha_1$ and $\alpha_2$ are 0, and equation (2.11) simplifies to $\lambda_{i,0}$, which is equivalent to the inverse logit of $g_i$ in the DINO model. If only attribute 1 has been mastered, the $\alpha_2$ will be equal to 0, and equation (2.11) simplifies to $\lambda_{i,0} + \lambda_i$, which is equivalent to $(1 - s_i)$ in the DINO model. Finally, if both attributes have been mastered, then equation (2.11) becomes $\lambda_{i,0} + \lambda_i (1 + 1 - 1 \times 1) = \lambda_{i,0} + \lambda_i$, which is identical the result when only one attribute was mastered.

Because the LCDM is able to encompass this variety of DCMs, the choice of compensatory versus non-compensatory DCM becomes irrelevant. Instead, the saturated LCDM can be estimated, and if the items truly follow the DINA, DINO, or other lower-level DCM, the estimated
parameters will reflect this. Further the LCDM provides a framework for testing the assumptions of these other DCMs. For example, two models could be fit to the same data: one fully saturated LCDM, the other with constraints on the item parameters. A likelihood ratio test can then be performed to determine if the reduced model fits as well as the saturated model.

### 2.4 Structural models

To this point, the discussion has focused on the measurement model of DCMs. Recall from equation (2.1), reprinted here, that this is only one piece of the diagnostic model.

\[
P(X_r = x_r) = \sum_{c=1}^{C} \nu_c \prod_{i=1}^{I} \pi_{ic}^{x_{ir}} (1 - \pi_{ic})^{1-x_{ir}}
\]

(2.12)

The measurement model, which relates the attributes to the observed item responses, is concerned with estimated \( \pi_{ic} \). The structural model is focused on \( \nu_c \). In DCMs, \( \nu_c \) represents the base rate probability of each latent class (Rupp et al., 2010). The base probabilities allow for the calculation of mastery rates for each attribute marginally, as well as the correlations between the attributes. In an assessment with \( A \) attributes there are \( 2^A \) latent classes. Because all elements of \( \nu \) must sum to 0, there are \( 2^A - 1 \) parameters to estimate, as the final element can be calculated by taking 1 minus the sum of the other elements.

Estimating each of these probabilities directly is referred to as the “unstructured” or “unconstrained” structural model (Rupp et al., 2010). By estimating the probabilities directly, it is possible to observe if there are any classes that have few respondents, possibly indicating the presence of an attribute hierarchy as described by (Templin & Bradshaw, 2014a). However, this type of unconstrained model can cause problems in high dimensional attribute spaces. Because the number of structural parameters to be estimated is \( 2^A - 1 \), the number of parameters increases exponentially with each added attribute. For example, a five attribute assessment requires \( 2^5 - 1 = 31 \) parameters, whereas a 10 attribute assessment would require \( 2^{10} - 1 = 1,023 \) parameters. Thus, it is often desirable to reduce the number of parameters that need to be estimated.
Two such approaches for reducing the structural model are the unstructured tetrachoric model (Hartz, 2002) and structured tetrachoric model (de la Torre & Douglas, 2004). These approaches work well in many instances (e.g., if the primary interest is the relationships between the attributes), however, they can be rather restrictive in what can be estimated. For example, suppose that and unstructured tetrachoric model is utilized, and it is determined that the structural model has been reduced too much. With this method, it is not a straightforward proposition as to how to add parameters back in without going all the way back to the unconstrained model. Additionally, suppose that a researcher is interested in both the potential hierarchical structure of attributes and the attributes’ associations. How should the researcher reduce the model, and achieve both of these desired outcomes? Rupp et al. (2010) suggest a “top-down approach” using log-linear models, which is described in the following section (section 2.4.1).

2.4.1 Log-linear structural models

The log-linear approach to structural models was proposed by (Henson & Templin, 2005), and is very similar to the log-linear model that was used to define the measurement model of the LCDM in section 2.3.1. Specifically, the kernel for latent class \( c \) can be defined as follows:

\[
\text{kernel}_c = \sum_{a=1}^{A} \gamma_{1,(a)} \alpha_{ca} + \sum_{a=1}^{A-1} \sum_{a'=a+1}^{A} \gamma_{2,(a,a')} \alpha_{ca} \alpha_{ca'} + \ldots + \sum_{a=1}^{A} \prod_{a=1}^{A} \alpha_{ca} \tag{2.13}
\]

The parameters included for latent class \( c \) are a main effect for each attribute that has been mastered by individuals in the class, as well interactions between the mastered attributes (two-way up to \( A \)-way, where \( A \) is the total number of attributes assessed). For an assessment measuring two attributes, the structural model would be defined as outlined in Table 2.3.

The fully saturated log-linear structural model is equivalent to the unconstrained structural model. However, this parameterization has many benefits over the tetrachoric methods. First, using this method allows for a hypothesis test on each of the estimated parameters. Thus, non-significant parameters can be removed from the model, allowing for model reduction to occur.
Table 2.3: Log-linear structural model for a 2-attribute assessment

<table>
<thead>
<tr>
<th>Class</th>
<th>Attribute Profile</th>
<th>Kernel</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>[0,0]</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>[1,0]</td>
<td>γ₁(1)</td>
</tr>
<tr>
<td>3</td>
<td>[0,1]</td>
<td>γ₁(2)</td>
</tr>
<tr>
<td>4</td>
<td>[1,1]</td>
<td>γ₁(1) + γ₁(2) + γ₂(1,2)</td>
</tr>
</tbody>
</table>

without enforcing a less flexible structure. This parameterization can also be used to reduce the structural model prior to estimation. For example, Xu & von Davier (2008) used a log-linear structural model in their analysis of data from the National Assessment of Educational Progress. In the structural model, they allow for only main effects, two-way and three-way interactions. Estimating only main effects results in independent attributes. The addition of the two-way interaction allows for variances (and therefore correlations) to be estimated. Finally, the three-way interaction allows for the third moment, skewness, to be captured. Although the log-linear structural model may not be as intuitive as the tetrachoric models to those familiar with structural models in structural equation modeling and multidimensional item response theory, its flexibility makes it easier to remove parameters from the structural model when the attribute structure is unclear a priori.

2.5 Model reduction

As has been generally discussed thus far, model reduction is the process of removing parameters from the model in order to create a more parsimonious and efficient model (Templin & Bradshaw, 2014b), while still maintaining a structure that is capable of capturing the complexity of the data. The parameters can be removed from either the structural model or the measurement model. In practice, model reduction can take place in different contexts. The first is what usually comes to mind when thinking of model reduction. That is, removing parameters after the initial estimation of the model. However, model reduction is also common when the initial model fails to converge. In this scenario, without the parameter estimates and hypothesis tests from the initial model, heuristic decisions must be made in order to reduce the model to a structure that is estimable. Although
different, understanding these processes is critical to the interpretation of the results. Before examining prior research on this using DCMs, the related literature from other latent variable models such as structural equation modeling and item response theory will be examined.

2.5.1 Model reduction in structural equation modeling

As in DCMs, in structural equation modeling, the measurement model relates the observed variables to the latent traits and the structural model defines the relationships between the latent traits. As such, both parts of the structural equation model can be reduced. In practice, this is a multi-stage process. Both Kline (2002) and Ullman (2012) suggest first fitting each measurement model separately. In other words, for each latent variable, fit a unidimensional model first. Then, add or remove parameters as necessary in order to ensure model fit. Once all of the unidimensional models have been assessed for model fit, they can be estimated simultaneously in the structural equation model, and the structural model can be reparameterized as needed to ensure the fit of the whole model.

Thus, the structural equation modeling world seems to follow a measurement model first, then structural model approach to model reduction. In an examination of the relationships between mental toughness, motivation, and emotion in sports, Perry, Nicholls, Clough, & Crust (2015) examined the factors from each questionnaire separately before combining them into the full model. They found that the full model had significantly better fit when the measurement models were adjusted prior to the estimation of the full model. Similarly, Burkholder & Harlow (2003) followed this procedure to reduce their model investigating HIV behavior risk. In this model, there was no reduction of the measurement model as each factor was just-identified. Thus, there was only reduction at the structural level, where all non-significant regression coefficients were removed.

However, it should be noted that model modifications are not limited to the removal of parameters in structural equation models. It is also common to use modification indices to locate places in the model where there is misfit and add parameters to improve the overall fit (Brown, 2006; Kaplan, 2009). These parameters could include additional regression paths, covariances be-
between latent factors, or residual covariances. Thus, when structural equation models are modified following the initial estimation it is common practice to not only reduce the model by removing non-significant parameters, but also add additional parameters in order to ensure model fit.

2.5.2 Model reduction in item response theory

Unlike structural equation modeling, model reduction is relatively uncommon in item response theory. There are a few possible reasons for this. First, the majority of operationally used item response theory models are unidimensional, with multidimensional item response theory models having yet to see wide spread operational use (see Fukuhara & Kamata, 2011; Reckase, 1997; Sinharay, 2010; Thissen & Steinberg, 1986). In unidimensional models, there are no relationships between latent variables to estimate, as there is only one. Thus, only the measurement model is of consequence. In unidimensional item response theory models, this comes down to the selection of the number of parameters to be included (i.e., 1-parameter logistic model, 2-parameter logistic model, or 3-parameter model). Thus, after choosing an initial model, the options are to either reduce the model by removing items that don’t fit, or change models to add additional parameters.

In contrast, multidimensional item response theory models do offer opportunities for model reduction. In multidimensional models, the covariance structure of the latent factors can be reduced, as well as respecifying the latent traits that are measured by each item. However, although there are a few examples of various specifications being tested (see Kingston & McKinley, 1988; McKinley & Kingston, 1988), this is typically not done in practice. Indeed, there is no mention at all of modifying the structure of the multidimensional model following estimation in the most widely cited textbook on multidimensional item response theory models (see Reckase, 2009). Instead, a saturated covariance matrix is estimated for the latent traits, and decisions about the measurement model parameters are confined to removing items that don’t fit the model.
2.5.3 Model reduction in DCMs

In diagnostic models, the model reduction process is more similar to structural equation modeling than item response theory, in that it is a common practice to reduce the model by removing non-significant parameters after the estimation of the initial model. For example, Jurich & Bradshaw (2013) used the LCDM with a log-linear structural model to scale the Socialcultural Dimension Assessment version 6 (Halonen, Harris, Pastor, Abrahamson, & Huffman, 2005). In the estimation of the LCDM, four separate structural models were defined *a priori*: the fully saturated log-linear model, reduced model where three- and four-way interactions were removed (constrained to be 0), and a model with only main effects and two-way interactions that were constrained to be equal. Initially, Jurich & Bradshaw (2013) estimated the model with a fully saturated measurement and structural model. Following this initial estimation they removed non-significant parameters from the measurement model, before using the reduced measurement model to evaluate the structural models.

A similar approach was used by Bradshaw, Izsák, Templin, & Jacobson (2014) in their analysis of the Diagnosing Teachers’ Multiplicative Reasoning assessment. In this analysis, a fully saturated LCDM and log-linear structural model were used in the initial estimation. However, when three- and four-way interaction terms were specified in the structural model, the model was not able to converge. Thus, they settled on a reduced structural model with only main effects and two-way interactions included. Using the reduced structural model, Bradshaw et al. proceeded to remove non-significant terms from the measurement model. This same procedure was followed by de la Torre, Ark, & Rossi (2015) in their analysis of the Dutch version of the Millon Clinical Multiaxial Inventory-III (T. Millon, Millon, Davis, & Grossman, 2009). Using a saturated structural model, the measurement model was reduced by removing non-significant terms from each item. However, no further model reduction was done to the structural model.

As can be seen from these studies, there is an inconsistency as to the order in which model reduction should occur in DCMs. Bradshaw et al. (2014) reduced the structural model first out of necessity due to convergence, whereas Jurich & Bradshaw (2013) elected to reduce the mea-
surement model first. However, this has not, to this point, been an investigation as to which of the processes should be preferred.

One option would be to follow the direction of the structural equation modeling literature and always reduce the measurement model first. This is potentially problematic for a few reasons. First, structural equation models assume that the data is normally distributed, whereas diagnostic models assume dichotomous items. It is possible to have non-dichotomous data (e.g., Skrondal & Rabe-Hesketh, 2004), but that is beyond the scope of this study. Additionally the latent variables in structural equation models are generally continuous, whereas DCMs use categorical latent variables. Because the observed data and latent variable space both follow different distributions, it’s possible that the best practices may differ between the two models.

Additionally, there is the added complexity of precisely how the measurement model is estimated in structural equation models. Recall from section 2.5.1 that the recommended practice to estimate each latent variable’s own measurement model first in order to ensure fit (Kline, 2002). However the measurement model for DCMs require multiple attributes in order to estimate the interaction effects (see equation (2.8)). Thus, the measurement models could only be estimated separately when there was a simple specification of the Q-matrix where each item measured only one attribute. Therefore, the approach taken by the structural equation modeling community is unlikely to be feasible for most DCM assessments.

2.6 The current study

Despite the uncertainty in the process of model reduction, this is a crucial aspect of DCM estimation. Rojas, de la Torre, & Olea (2012) showed that estimating high level interaction terms when unnecessary can decrease the classification accuracy compared to a reduced model. However, Rojas et al. (2012) only examined reduction of the measurement model. Therefore, the present study seeks to examine reduction of both the measurement and structural models to answer the following research questions:
1. Does choosing different a model reduction process impact the output of the model?

2. What are the benefits of reducing the measurement and/or structural model(s)?

3. Are there advantages or disadvantages to the order of reduction (i.e., measurement or structural reduction first)?

This is accomplished through two studies. The first is a pilot study to further demonstrate the importance of the order of model reduction. Specifically the Diagnosing Teachers’ Multiplicative Reasoning assessment data (as described in Bradshaw et al., 2014) is reduced in different orders to compare the values of the resulting parameter estimates. Second, given the results of the pilot study, and Monte Carlo simulation study is conducted to examine the effects of model reduction and the order of reduction under a variety of data generation and dimensionality conditions.
Chapter 3

Pilot Study

In order to better assess the impact of the order of model reduction, a pilot study was conducted on the Diagnosing Teachers’ Multiplicative Reasoning (DTMR) assessment data, as described in Bradshaw et al. (2014). In this study, the DTMR data set was analyzed using a variety of model reduction processes to determine if the selected process has an impact on the resulting model parameters. Thus, the pilot study was designed to answer the first research question defined in section 2.6. This is an important first step as without evidence that the model reduction process has an impact on the results, there would be little motivation for a more thorough analysis via simulation.

3.1 Method

3.1.1 DTMR data

The DTMR assessment consists of 28 dichotomously scored items that together measure 4 attributes related to educators’ understanding of multiplicative reasoning (Bradshaw et al., 2014):

1. Referent Units (RU): recognizing which whole the fraction refers to,
2. Partitioning and Iterating (PI): splitting a whole into equal pieces repeatedly to achieve larger fractions,
3. Appropriateness (APP): determining the correct mathematical operation from a problem statement, and
4. Multiplicative Comparison (MC): evaluating the ratio of one value to another.
Of the 28 items, Referent Units is measured by 16, Partitioning and Iterating by 10, Appropriateness by 6, and Multiplicative Comparison by 10. This totals adds to more than 28 because several items measure more than one attribute. Specifically, there are 14 items measuring only 1 attribute and 14 that measure 2 attributes. The complete saturated Q-matrix for the DTMR assessment can be see in Table 3.1. For example, item 1 measures only the Referent Units attribute, whereas item 5 measures both the Referent Units and Multiplicative Comparison attributes.

Table 3.1: DTMR Q-matrix

<table>
<thead>
<tr>
<th>Item</th>
<th>Item Name</th>
<th>RU</th>
<th>PI</th>
<th>APP</th>
<th>MC</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>3</td>
<td>3</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>4</td>
<td>4</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>5</td>
<td>5</td>
<td>1</td>
<td>0</td>
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</table>
In total 990 math teachers took the assessment. Bradshaw et al. (2014) reported that sample demographics were consistent with a representative national sample. For a complete description of the sample characteristics and data collection process, see the original description of the DTMR in Bradshaw et al. (2014).

### 3.1.2 Model estimation

In order to assess the impact of various model reduction processes, the LCDM was estimated for the DTRM using five different reduction methods:

1. Simultaneous reduction: after estimating the fully saturated model, all non-significant parameters from both the measurement and the structural models are removed and the model is re-estimated,
2. Measurement reduction: after estimating the fully saturated model, all non-significant parameters from the measurement model only are removed and the model is re-estimated,
3. Structural reduction: after estimating the fully saturated model, all non-significant parameters from the structural model only are removed and the model is re-estimated,
4. Measurement-Structural reduction: after estimating the measurement reduction model, all non-significant parameters from the structural model only are removed and the model is re-estimated, and
5. Structural-Measurement reduction: after estimating the structural reduction model, all non-significant parameters from the measurement model only are removed and the model is re-estimated.

The different ordering processes for model reduction, and their relationships to each other, are represented visually in Figure 3.1. The significance of each parameter was determined by the p-value derived from the Wald test provided by Mplus. This test provides a p-value for the null hypothesis that the parameter is equal to zero. Parameters with a p-value greater than 0.05 were determined to be non-significant, and therefore removed at the corresponding stage of model estimation.
reduction. It should be noted that in the model reduction processes outlined, a constraint was put in place to prevent the removal of item intercepts. An intercept must be defined in order to ensure that all respondents have an estimated probability of a correct response. Thus, item intercepts remained in the model, regardless of their significance.

![Flowchart of model reduction processes](image)

**Figure 3.1: Flowchart of model reduction processes**

In practical terms, a non-significant intercept means that the log-odds of a respondent who hasn’t mastered any of the required attributes is not significantly different from 0 (this corresponds to a probability of 0.5). Thus, these cases may represent easier, or highly guessable items, where non-masters of the required traits still have a relatively high probability of success. In contrast, non-significant main effects and interactions represent instances where the increase in the log-odds for masters of the given attribute (or combination of attributes in the case of interactions) is not significantly different from 0. Thus, after removing these parameters, the affected respondents would have the same probability of providing a correct response as respondents who had not mastered the required attributes. In the extreme case where all parameters for an item are removed
except for the intercept, all respondents would have the same probability of success, regardless of
their profile of attribute mastery.

All analyses for the pilot study were conducted in *Mplus* version 7.4 (L. K. Muthén & Muthén,
1998) via the *MplusAutomation* package (Hallquist & Wiley, 2018) in *R* version 3.4.3 (R Core
Team, 2017). *Mplus* code for the estimation of the LCDM was generated in *R* using custom scripts

### 3.2 Results

The final estimate and associated standard error for each parameter from each of the model reduc-
tion processes are presented in their own tables in order to easily compare across model reduction
processes. Table 3.2, Table 3.3, Table 3.4, Table 3.5, Table 3.6, Table 3.7, Table 3.8, Table 3.9,
and Table 3.10 show the results of the measurement model parameters, and Table 3.11 shows the
results of the structural model parameters.

#### 3.2.1 Measurement model results

Although it is tempting to compare the point estimates and standard errors for the parameters
across model reduction techniques, this is ill-advised. Because the main effects and interactions
are conditional on other parameters in the model, the exclusion of parameters will change the
interpretation of the other parameters. Thus, changes in point estimates and standard errors may
be expected. Thus, these results are most useful for comparing which parameters are ultimately
retained in the model. In general we can see that the choice of the model reduction method can
have a profound impact on which parameters are ultimately retained in the model. The exception
to this is the item intercepts (Table 3.2). Because model reduction was constrained to never remove
item intercepts, all intercepts are estimated, no matter which reduction technique was used. This is
not the case for main effects. For example, when examining the main effects for attribute 1 (Table
3.3), the main effect for item 13 was only retained when either the structural model only was
Table 3.2: DTMR estimates of item intercepts, $\lambda_{i,0}$

<table>
<thead>
<tr>
<th>Item</th>
<th>Simultaneous Reduction</th>
<th>Measurement Reduction</th>
<th>Structural Reduction</th>
<th>Measurement-Structural Reduction</th>
<th>Structural-Measurement Reduction</th>
</tr>
</thead>
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</tr>
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* Parentheses show the standard error of the estimate.

Reduced, or when the structural model was reduced first. This is because when the measurement model was reduced first, this main effect was non-significant, and thus reduced out of the model. However, reduction of the structural model first changed the parameter estimate and associated p-value enough for the parameter to be significant, and thus retained.
Table 3.3: DTMR estimates of item main effects for Referent Units, $\lambda_{i,1,(1)}$

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</table>

* Parentheses show the standard error of the estimate.
† Missing values indicate the parameter was removed in the reduction process.

Table 3.4: DTMR estimates of item main effects for Partitioning and Iterating, $\lambda_{i,1,(2)}$

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<th>Structural Reduction</th>
<th>Measurement-Structural Reduction</th>
<th>Structural-Measurement Reduction</th>
</tr>
</thead>
<tbody>
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</table>

* Parentheses show the standard error of the estimate.
† Missing values indicate the parameter was removed in the reduction process.
Table 3.5: DTMR estimates of item main effects for Appropriateness, $\lambda_{i,1,(3)}$

<table>
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<tr>
<th>Item</th>
<th>Simultaneous Reduction</th>
<th>Measurement Reduction</th>
<th>Structural Reduction</th>
<th>Measurement-Structural Reduction</th>
<th>Structural-Measurement Reduction</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>1.30 (0.20)</td>
<td>1.27 (0.06)</td>
<td>1.30 (0.22)</td>
<td>1.22 (0.24)</td>
<td>1.30 (0.21)</td>
</tr>
<tr>
<td>8</td>
<td>3.63 (0.34)</td>
<td>4.32 (0.11)</td>
<td>4.20 (0.58)</td>
<td>4.55 (0.68)</td>
<td>3.83 (0.51)</td>
</tr>
<tr>
<td>9</td>
<td>2.02 (0.21)</td>
<td>2.16 (0.14)</td>
<td>2.17 (0.25)</td>
<td>2.24 (0.23)</td>
<td>2.09 (0.25)</td>
</tr>
<tr>
<td>10</td>
<td>0.84 (0.18)</td>
<td>0.84 (0.04)</td>
<td>0.88 (0.18)</td>
<td>0.81 (0.18)</td>
<td>0.87 (0.18)</td>
</tr>
<tr>
<td>11</td>
<td>1.89 (0.20)</td>
<td>1.81 (0.19)</td>
<td>1.86 (0.24)</td>
<td>1.72 (0.21)</td>
<td>1.86 (0.21)</td>
</tr>
<tr>
<td>26</td>
<td>0.00 (0.00)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Parentheses show the standard error of the estimate.
† Missing values indicate the parameter was removed in the reduction process.

Table 3.6: DTMR estimates of item main effects for Multiplicative Comparison, $\lambda_{i,1,(4)}$

<table>
<thead>
<tr>
<th>Item</th>
<th>Simultaneous Reduction</th>
<th>Measurement Reduction</th>
<th>Structural Reduction</th>
<th>Measurement-Structural Reduction</th>
<th>Structural-Measurement Reduction</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td></td>
<td></td>
<td>0.27 (0.31)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8</td>
<td></td>
<td></td>
<td>0.54 (0.50)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>13</td>
<td>4.87 (0.60)</td>
<td>4.89 (2.00)</td>
<td>5.19 (2.08)</td>
<td>4.87 (0.60)</td>
<td>4.47 (0.56)</td>
</tr>
<tr>
<td>14</td>
<td>4.13 (0.66)</td>
<td>4.28 (0.64)</td>
<td>3.80 (0.84)</td>
<td>4.12 (0.65)</td>
<td>4.21 (0.81)</td>
</tr>
<tr>
<td>15</td>
<td>4.66 (0.91)</td>
<td>4.67 (0.46)</td>
<td>4.19 (0.81)</td>
<td>4.75 (0.96)</td>
<td>4.44 (0.91)</td>
</tr>
<tr>
<td>16</td>
<td></td>
<td></td>
<td>0.25 (0.25)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>18</td>
<td></td>
<td></td>
<td>0.48 (0.28)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>20</td>
<td>1.25 (0.50)</td>
<td></td>
<td>1.22 (0.28)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>21</td>
<td>0.63 (0.36)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>22</td>
<td>0.83 (0.34)</td>
<td></td>
<td>0.52 (0.27)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Parentheses show the standard error of the estimate.
† Missing values indicate the parameter was removed in the reduction process.

A similar, but more extreme pattern can be seen in the interactions in the measurement model (Tables 3.7, 3.8, 3.9, 3.10). Interactions between the attributes were only retained when the structural model alone was reduced (in which case all interactions from the measurement model were kept) or when the structural model was reduced first (in which case some of the interactions were retained). If the measurement model was reduced before the structural model, or with the structural model simultaneously, interaction terms from the measurement model were never retained.

Finally, as mentioned above, it is generally ill-advised to compare point estimates and standard
Table 3.7: DTMR estimates of item interactions between Referent Units and Partitioning and Iterating, $\lambda_{i,2,(1,2)}$

<table>
<thead>
<tr>
<th>Item</th>
<th>Simultaneous Reduction</th>
<th>Measurement Reduction</th>
<th>Structural Reduction</th>
<th>Measurement-Structural Reduction</th>
<th>Structural-Measurement Reduction</th>
</tr>
</thead>
<tbody>
<tr>
<td>19</td>
<td>1.35 (1.95)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>24</td>
<td>0.91 (0.43)</td>
<td></td>
<td></td>
<td>1.04 (0.27)</td>
<td></td>
</tr>
<tr>
<td>25</td>
<td>0.04 (0.84)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>28</td>
<td>-1.36 (0.94)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
* Parentheses show the standard error of the estimate.
† Missing values indicate the parameter was removed in the reduction process.

Table 3.8: DTMR estimates of item interactions between Referent Units and Multiplicative Comparison, $\lambda_{i,2,(1,4)}$

<table>
<thead>
<tr>
<th>Item</th>
<th>Simultaneous Reduction</th>
<th>Measurement Reduction</th>
<th>Structural Reduction</th>
<th>Measurement-Structural Reduction</th>
<th>Structural-Measurement Reduction</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>-0.16 (0.88)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>13</td>
<td>3.65 (6.32)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>14</td>
<td>-1.93 (1.57)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>15</td>
<td>-2.45 (1.14)</td>
<td></td>
<td></td>
<td>1.32 (0.28)</td>
<td></td>
</tr>
<tr>
<td>16</td>
<td>-0.25 (0.25)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
* Parentheses show the standard error of the estimate.
† Missing values indicate the parameter was removed in the reduction process.

Table 3.9: DTMR estimates of item interactions between Partitioning and Iterating and Multiplicative Comparison, $\lambda_{i,2,(2,4)}$

<table>
<thead>
<tr>
<th>Item</th>
<th>Simultaneous Reduction</th>
<th>Measurement Reduction</th>
<th>Structural Reduction</th>
<th>Measurement-Structural Reduction</th>
<th>Structural-Measurement Reduction</th>
</tr>
</thead>
<tbody>
<tr>
<td>18</td>
<td>-0.04 (0.47)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>20</td>
<td>0.79 (0.71)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>21</td>
<td>1.95 (2.27)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>22</td>
<td>1.54 (1.46)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
* Parentheses show the standard error of the estimate.
† Missing values indicate the parameter was removed in the reduction process.

errors across model reduction methods. The exception to this rule is the item intercepts, because these parameters are not dependent on the other terms in the model (i.e., the intercept always
Table 3.10: DTMR estimates of item interactions between Appropriateness and Multiplicative Comparison, $\lambda_{i,2,(3,4)}$

<table>
<thead>
<tr>
<th>Item</th>
<th>Simultaneous Reduction</th>
<th>Measurement Reduction</th>
<th>Structural Reduction</th>
<th>Measurement-Structural Reduction</th>
<th>Structural-Measurement Reduction</th>
</tr>
</thead>
<tbody>
<tr>
<td>8</td>
<td>-0.54 (0.50)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Parentheses show the standard error of the estimate.
† Missing values indicate the parameter was removed in the reduction process.

represents the log-odds of providing a correct response when none of the attributes have been mastered). Thus, it is possible to compare these parameters across model reductions methods. It is clear that although these parameters are generally similar, the standard errors sometimes vary wildly (for example, Table 3.2). For instance, the standard error of the intercept on item four is consistently around 0.10, except for when only the measurement model is reduced. In this situation, the standard error is 14.55. A similar pattern can be seen in the intercepts for items 5, 18, 19, and 23.

### 3.2.2 Structural model results

The parameters of the structural model also show variability between model reduction processes. As with the measurement model parameters, different model reduction processes result in different parameters ultimately being retained in the model. Table 3.11 shows, for example, that 3-way interactions are almost always removed unless the measurement model is reduced first. The exception is the 3-way interaction between the Referent Units, Appropriateness, and Multiplicative Comparison attributes. Notably, this interaction appears to be relatively unstable across model reduction process, varying from 2.40 to 14.94. However, this variability in point estimates should be interpreted with caution. As was the case when examining the measurement model parameters, the structural parameters are also conditional upon other parameters, and thus, fluctuations may be expected depending on the inclusion or exclusion of the other parameters.

The variation in which structural parameters are ultimately retained in the model can have
serious implications for classification of respondents. Given that these parameters govern the base rate probabilities of membership in each of the attribute profiles, these differences in included parameters could lead to differences in the classification of respondents.

Table 3.11: DTMR estimates of structural parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Simultaneous Reduction</th>
<th>Measurement Reduction</th>
<th>Structural Reduction</th>
<th>Measurement-Structural Reduction</th>
<th>Structural-Measurement Reduction</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\gamma_{1,(1)}$</td>
<td>-5.11 (0.01)</td>
<td>-3.49 (1.58)</td>
<td>-10.06 (0.43)</td>
<td>-2.93 (0.62)</td>
<td>-6.85 (2.28)</td>
</tr>
<tr>
<td>$\gamma_{1,(2)}$</td>
<td>-2.10 (0.04)</td>
<td>-2.92 (1.77)</td>
<td>-1.65 (0.34)</td>
<td>-1.21 (0.24)</td>
<td>-1.80 (0.34)</td>
</tr>
<tr>
<td>$\gamma_{1,(3)}$</td>
<td>-1.54 (0.05)</td>
<td>-1.28 (0.35)</td>
<td>-1.40 (0.37)</td>
<td>-0.75 (0.33)</td>
<td>-1.55 (0.37)</td>
</tr>
<tr>
<td>$\gamma_{1,(4)}$</td>
<td>-1.06 (0.06)</td>
<td>-1.24 (0.13)</td>
<td>-0.84 (0.28)</td>
<td>-0.95 (0.24)</td>
<td>-0.91 (0.28)</td>
</tr>
<tr>
<td>$\gamma_{2,(1,2)}$</td>
<td>3.44 (0.01)</td>
<td>-2.96 (0.04)</td>
<td>8.75 (0.52)</td>
<td>-8.35 (0.66)</td>
<td>5.56 (2.24)</td>
</tr>
<tr>
<td>$\gamma_{2,(1,3)}$</td>
<td>0.53 (0.02)</td>
<td>-8.16 (3.62)</td>
<td>1.53 (0.78)</td>
<td>-6.93 (0.68)</td>
<td>0.36 (0.93)</td>
</tr>
<tr>
<td>$\gamma_{2,(1,4)}$</td>
<td>-5.24 (3.57)</td>
<td></td>
<td>1.53 (0.78)</td>
<td>-6.93 (0.68)</td>
<td>0.36 (0.93)</td>
</tr>
<tr>
<td>$\gamma_{2,(2,3)}$</td>
<td>2.23 (0.02)</td>
<td>2.46 (1.44)</td>
<td>1.87 (0.46)</td>
<td>2.02 (0.42)</td>
<td></td>
</tr>
<tr>
<td>$\gamma_{2,(2,4)}$</td>
<td>2.31 (1.34)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\gamma_{2,(3,4)}$</td>
<td>1.89 (0.02)</td>
<td>1.70 (0.34)</td>
<td>1.51 (0.45)</td>
<td>1.41 (0.36)</td>
<td>1.81 (0.38)</td>
</tr>
<tr>
<td>$\gamma_{3,(1,2,3)}$</td>
<td>13.91 (3.71)</td>
<td></td>
<td></td>
<td></td>
<td>18.28 (0.78)</td>
</tr>
<tr>
<td>$\gamma_{3,(1,2,4)}$</td>
<td>10.38 (3.60)</td>
<td></td>
<td></td>
<td></td>
<td>10.79 (0.66)</td>
</tr>
<tr>
<td>$\gamma_{3,(1,3,4)}$</td>
<td>2.40 (0.03)</td>
<td>14.94 (0.12)</td>
<td>7.15 (0.87)</td>
<td>7.85 (0.70)</td>
<td>4.73 (2.38)</td>
</tr>
<tr>
<td>$\gamma_{3,(2,3,4)}$</td>
<td>-1.65 (1.62)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$\gamma_{4,(1,2,3,4)}$</td>
<td>-0.49 (0.03)</td>
<td>-18.66 (1.67)</td>
<td>-6.98 (0.68)</td>
<td>-16.70 (1.04)</td>
<td>-3.18 (2.34)</td>
</tr>
</tbody>
</table>

* Parentheses show the standard error of the estimate.
† Missing values indicate the parameter was removed in the reduction process.

Figure 3.2 shows the change in the number of respondents classified in each attribute profile under each model reduction processes. Although the counts are fairly consistent across most attribute profiles, the counts for profiles 15 ([0,1,1,1]) and 16 ([1,1,1,1]) do show large discrepancies. There are about 50 respondents classified as master of Partitioning and Iterating, Appropriateness, and Multiplicative Comparison under the measurement-structural reduction process, but around 150 when using all other processes. The attribute profile where all attributes is mastered ([1,1,1,1]) is even more volatile, with the number of respondents placed into this class varying from 200 to nearly 350 depending on the reduction method.
3.3 Conclusions

The results from the pilot study provide evidence that the choice of model reduction process can significantly influence not only the estimates of the parameters, but also the classification of respondents into attribute profiles. However, without knowing the true values of parameters and attribute profiles of the respondents, it is impossible to know which model reduction process should be chosen. Therefore, a simulation study will be conducted in which the true values of the parameters and attribute profiles are known. In this way, it will be possible to measure how accurately each
model reduction process is able to recover parameter values and respondent attribute profiles. The results of the simulation will then be able to inform which method is best suited to analyze the DTMR data.
Chapter 4

Methods

The current study investigates the performance of the different model reduction processes using a Monte Carlo simulation. The simulation will be designed to follow the structure and characteristics of the DTMR data and the corresponding estimated parameters.

4.1 Overview of Monte Carlo methods

Broadly speaking, Monte Carlo simulations can be used to evaluate the performance of different or competing methodologies (Rubinstein & Kroese, 2017). As discussed in section 3.3, when using real data it is difficult to evaluate competing procedures, because the true values of the parameters are unknown. Thus, the estimates from different procedures can be compared to each other, but cannot be compared to what the real value is. With Monte Carlo methods, data is generated from a model with known parameters. The different procedures can be used on the simulated data, and then the resulting estimates compared to the actual values that were used to generate the data.

The general process of a Monte Carlo simulation is as follows:

1. Define distributions for model parameters,
2. Randomly draw model parameters from previously defined distributions,
3. Simulate response data from the drawn model parameters,
4. Estimate the model using simulated response data, and
5. Compare estimated parameters to the parameters that were randomly drawn and used to simulate the data.
Monte Carlo studies have been used extensively in psychometric research (see Feinberg & Rubright, 2016 for a survey of simulation studies in educational measurement). There are disadvantages to this methodology however (e.g., Harwell, Stone, Hsu, & Kirisci, 1996). In the simulation of data, only parameters that are defined influence the generation of response data. Thus, if the model specified in the simulation study is inconsistent with reality, the simulation may not be informative. Additionally, the data generated will be perfectly model fitting (unless misfit is introduced in the data generation process). This means that the findings of the simulation study may not hold when there is misfit in the data. In other words, the main disadvantage of Monte Carlo studies is that simulation environment may not generalize to empirical data. Therefore, when designing a Monte Carlo study it is important to make the conditions and parameter distributions as realistic as possible (Feinberg & Rubright, 2016).

4.2 The current simulation

In order to make this simulation as realistic as possible, it will be modeled after the DTMR assessment and the results from the pilot study.

4.2.1 Simulation conditions

The DTMR assessment consists of 28 items measuring four attributes. The simulation study will use three and four attributes in order to assess the effects of increasingly complex structural models. Additionally, each condition will hold the number of items constant at 30 to reduce the number of conditions in the simulation. Three sample size conditions will be used: 500, 1000, and 5000. These sizes are representative of the DTMR sample size of 990, a smaller sample, and a large sample that would remove any sample size concerns for estimation. This will allow for the evaluation of sample size effects. Thus, there are two attribute conditions times three sample size conditions = six data generation conditions.

For each generated data set, three Q-matrices will be used to estimate the model: the true Q-
matrix, a 10 percent over-specified Q-matrix, and a 20 percent over-specified Q-matrix. In the over-specified Q-matrices, each value of zero in the Q-matrix will have either a 10 or 20 percent chance of being changed to a one, depending on the condition. Finally, for each Q-matrix, the model will be estimated in six ways:

1. Saturated model,
2. Simultaneous reduction,
3. Measurement reduction,
4. Structural reduction,
5. Measurement-Structural reduction, and

This results in a total of three Q-matrix conditions times six model estimation conditions = 18 estimation conditions. Therefore, there are six data generation conditions times 18 estimation conditions = 108 total conditions. Each condition will be replicated 100 times to ensure that robust estimates of the outcomes measures can be calculated for each condition.

4.2.2 Data generation process

When simulating data sets, the \( \gamma \) parameters that make up the structural model will be simulated first. There are no constraints on the values of \( \gamma \); however, based on the results of the pilot study, structural parameters will be drawn from a \( \mathcal{N}(0, 2) \) distribution. Using this distribution, most draws will come within 3 standard deviations of the mean, resulting in most values falling between -6 and 6. These values are consistent with what is seen in the DTMR pilot study (Table 3.11). Additionally, this range allows for large effects using the guidelines suggest by Chinn (2000), but also makes very large \( \gamma \) values unlikely.

Once the structural parameters have been generated, the base rate probabilities for each class can be calculated. These will in turn be used to generate attribute profiles for the simulated respondents. The attribute profiles, in conjunctions with item parameters, are used to calculate the
probability of each respondent answering each item correctly. The item intercepts will be drawn from a uniform distribution of [-5.00, 0.60]. This range is representative of the intercepts found in the pilot study, which range from -5.03 to 0.60 (Table 3.2), and therefore provides a realistic distribution of possible values. The main effects of the LCDM are constrained to be positive. Thus, these parameters will be drawn from of uniform distribution of (0.00, 5.00]. This is also based on the results on the pilot study, where the large estimated main effect was 5.19 (Table 3.3, Table 3.4, Table 3.5, and Table 3.6). Finally, the interaction terms are bounded on the lower end by \(-1 \times \text{smallest main effect}\). Thus the distribution for each interaction will be dependent on the main effects that are generated, but will all be bound on the upper end at 2.00. Only one interaction term from the pilot study fell beyond this range, the interaction between attributes 1 (RU) and 4 (MC) for item 13 in the structural reduction condition (Table 3.8). The item parameters and the attribute profiles are then used to calculate the probability of each respondent answering each item correctly. For each respondent/item combination then, a random uniform number is drawn between 0 and 1. If this random number is less than the probability of the respondent providing a correct response, a correct response is assigned (1), otherwise an incorrect response is assigned (0).

4.2.3 Model reduction process

Due to the potential issue in the convergence of DCMs noted in section 2.5.3, model reduction may occur in one of two ways for this study. The first method is used for the reduction of models that successfully converged. Given a converged model, parameters to remove are identified by their p-values, as was the case for analysis of the DTMR assessment (Chapter 3). The second method is for models that did not successfully converge. In this scenario, because p-values cannot be used to determine parameters that should be removed, all higher-order interaction parameters are flagged for removal (i.e., three-way interactions and greater). This heuristic is implemented in this study for two main reasons. First, it is highly unlikely that after collecting data, a researcher would simply give up on estimating the model if the fully saturated model did not converge. Thus, the
inclusion of this rule-based model reduction will inform practitioners on best practices. Secondly, the removal of higher-order interaction terms has already been implemented in the literature as a method for dealing with non-convergence (see Bradshaw et al., 2014 for an example). Accordingly, this implementation of this rule in the simulation study mimics the current practice of model estimation. A modified model reduction flow chart can be seen in Figure 4.1.

![Flowchart of simulation study reduction processes](image)

Figure 4.1: Flowchart of simulation study reduction processes

Note that in Figure 4.1, the rule-based model reductions can only occur after the fully saturated model. The logic behind this decision is as follows. If the fully saturated converges and the next model fails to converge, the practitioner should revert back to the model that successfully converged, rather than continuing to attempt to remove additional parameters. Conversely, if the fully saturated model fails to converge, and the next model also fails to converge, then the third model would result in the removal of both measurement and structural parameters using the heuristic. This would then be equivalent to the simultaneous reduction method using the heuristic, rendering third-step model redundant. Finally, regardless of the convergence of the fully saturated model, if
the second-step model converges, the model reduction process can continue to the third step using p-values.

### 4.2.4 Outcome measures

Results for models that were reduced from converged and non-converged models will be analyzed separately. This is to allow for the possibility that different methods of reducing the model may prefer different model reduction processes. Accordingly, convergence rates of each model reduction process will be an important outcome measure. In addition, estimated item and structural parameters will be compared to the true values. The bias and mean squared error of the estimates will be calculated to evaluate the ability of each model reduction process to recover the parameter values. In these calculations, parameters that were removed in the reduction process will have their estimates set to 0.

In addition to the recovery of model parameters, the recovery of attribute profiles will also be examined. This will be calculated at both the attribute and pattern level. This distinction differentiates how accurate attribute mastery was and how accurate profile assignment was. For example, assume a respondent had a true attribute profile of [1,0,1,0], and an estimated profile of [1,1,1,0]. At the pattern level, they were incorrectly assigned to an attribute profile. However, at the attribute level, 75% of attributes were correctly estimated. These measures will be evaluated using percent correct classification, and the quadratically weighted Cohen’s kappa (Cohen, 1960, 1968).

Another important outcome of the model fitting process is model data fit. Thus, it is important to investigate whether any reduction processes consistently provide better fit to the data. This will be assessed using the Akaike information criterion (AIC; Akaike, 1974), Bayesian information criterion (BIC; Schwarz, 1978), and the adjusted Bayesian information criterion (Sclove, 1987). These methods all measure the relative fit of a model compared to competing models. All the methods measure how well the model fits the data, with a penalty for the number of parameters. Within each measure, the model with the lowest index on the measure is the preferred model, taking into account both overall fit and model complexity.
Finally, diagnostics of the model reduction processes will be examined. Specifically, the rate at which each process results in the correct parameters being retained in the model will be examined. This will be examined separately for the measurement and structural models. Additionally, the distributions of the values and standard errors of parameters that are reduced with p-values will be evaluated. This will allow for an analysis of whether reduced parameters are being removed due to small parameter estimates or overly large standard errors.

4.2.5 Software

As with the pilot study, all models were estimated using *Mplus* version 7.4 (L. K. Muthén & Muthén, 1998) via the *MplusAutomation* package (Hallquist & Wiley, 2018) in *R* version 3.4.3 (R Core Team, 2017). *Mplus* code for the estimation of the LCDM was generated in *R* using custom scripts based on the work of Rupp & Wilhelm (2012) and Templin & Hoffman (2013). Data generation and evaluation of the models was carried out in *R* using the *dplyr* (Wickham, Francois, Henry, & Müller, 2018), *forcats* (Wickham, 2018a), *glue* (Hester, 2017), *lubridate* (Spinu, Grolemund, & Wickham, 2018), *portableParallelSeeds* (Johnson, 2016), *purrr* (Henry & Wickham, 2017), *readr* (Wickham, Hester, & Francois, 2017), *stringr* (Wickham, 2018b), *tibble* (Müller & Wickham, 2018), *tidyr* (Wickham & Henry, 2018), and *tidyselect* (Henry & Wickham, 2018) packages. All analyses were carried out on the Amazon Elastic Compute Cloud (EC2; Amazon Web Services, 2018).
Chapter 5

Results

Results from the simulation study are described in two sections. Section 5.1 summarizes the performance of the reduction processes when the parameters to reduce were determined by p-values from a converged model (i.e., p-value/p-value reduction). Section 5.2 summarizes results of the reduction processes for models that did not converge, and the parameters to reduce were determined by the higher order interaction heuristic (i.e., heuristic/p-value reduction).

Table 5.1 shows the convergence rates for the saturated model. As expected, the convergence rates decrease as the number of attributes increases, and decrease dramatically as the over specification of the Q-matrix increases. Interestingly, sample size had relatively little effect on the convergence rates of the saturated model, with rates staying fairly consistent with attribute and over specification conditions.

<table>
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5.1 Reduction by p-value

5.1.1 Convergence

When saturated model converged, reduction of measurement and structural parameters proceeded by using p-values to determine which parameters to reduce. Figure 5.1 shows the convergence rates for these reduction processes when the saturated model converged. Recall that measurement-structural and structural measurement reduction only occurred if the measurement and structural reductions, respectively, converged. Across all conditions, reduction processes where there measurement model was reduced first (i.e., simultaneous, measurement, and measurement-structural) tended to have higher convergence rates. This was most notable when the Q-matrix was over specified. In the most extreme case, the four-attribute condition with a 20 percent over specified Q-matrix, no reductions converged if the measurement model was not part of the initial reduction. However, because the saturated model very rarely converged for this condition (Table 5.1), the sample size is very limited.
5.1.2 Parameter recovery

To evaluate the performance of the reduction processes, the bias and mean square error of the parameter estimates can be examined. The bias represents the difference between the true value and the estimated value. The mean square error represents the average of the squared difference between the true and estimated values. Figure 5.2 and Figure 5.3 show the total bias and total mean square error, respectively, across all measurement model parameters when reducing using p-values. Because of some outlying data sets, biases with an absolute value greater than 100 and
mean square errors with a value greater than 100 have been excluded from the figures. Figures showing all biases and mean square errors, separated by the type of parameter can be seen in Appendix A.

Figure 5.2: Bias in measurement model main effect estimates when reducing using p-values
Figure 5.3: Mean square error in measurement model main effect estimates when reducing using p-values

Figure 5.2 shows that across all conditions there is relatively little bias in the measurement model parameters. This is especially true when the Q-matrix is correctly specified. The large negative bias seen in the measurement reduction condition for the three attribute and 10 percent over specified Q-matrix is due to a single three way interaction in one data set, as can be seen in Figure A.4. In contrast, there is relatively large mean square error values across conditions. As expected, this decreases as the sample size increases. Thus, these results suggest that larger sample sizes (i.e., greater than 1,000) are needed in order to ensure unbiased estimates of measurement
model parameters with low levels of error.

Figure 5.4 and Figure 5.5 show the bias and mean square error of the structural parameters respectively. Overall, there is very little bias and mean squared error in the structural parameters. The instances where larger bias and mean squared error are indicated (e.g., measurement reduction of a four attribute model with a 20% over specified Q-matrix) are instances where very few of the model actually converged. Thus, these values are based on only a few replications. Thus, these results suggests that given model convergence, the structural parameters are usually well estimated regardless of model reduction method.

Figure 5.4: Bias in structural model parameter estimates when reducing using p-values
Another critical measure of performance is the rate at which respondents are correctly classified as masters of the attributes. This is assessed at two levels: the individual attribute classifications, and the overall profile classification. As described in section 4.2.4, profile mastery and attribute mastery differentiate between two types of assignments that can occur. For example, a respondent may have a true profile of \([1,0,1,0]\) and an estimated profile of \([1,1,1,0]\). Here, the overall profile
assignment is incorrect (profile classification), but three out of the four individual attributes are correctly classified (attribute classification). Respondents were classified as a master of an attribute if their posterior probability of mastery was greater than or equal to .5, and non-master of the attribute otherwise. Figure 5.6 and Figure 5.7 show the attribute level agreement as measured by the average correct classification rate and average Cohen’s κ, respectively.

![Graph showing Attribute Correct Classification Rate](image)

Figure 5.6: Average correct classification rate of attribute mastery when reducing using p-values.
Across all conditions, both the correct classification rate and Cohen’s $\kappa$ show high rates of agreement between true and estimated attribute classifications. The exception is a four-attribute assessment with a sample size of only 500. Under these conditions, all model reduction processes showed lower rates of agreement than with larger samples or only three attributes.

The overall profile classification shows a similar pattern. Figure 5.8 and Figure 5.9 show the correct classification rate and Cohen’s $\kappa$ of the overall profiles. As expected, the overall profile agreement is consistently lower than the attribute level mastery classification agreement, although the classification agreement is generally consistent across all model reduction processes. Ad-
ditionally, as with the attribute classifications, profile classification agreement was lower in the four-attribute, 500 sample size conditions.

Figure 5.8: Average correct classification rate of profile assignment when reducing using p-values
### Figure 5.9: Average Cohen’s κ of profile assignment when reducing using p-values

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<td><img src="image8" alt="Graph" /></td>
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</tr>
</tbody>
</table>

#### 5.1.4 Model fit

Critical to the model selection process is model fit. Although different model reduction processes may all show adequate recovery of item parameters and respondent classification, these measures cannot provide insight as to whether or not the removal of the parameters significantly impacted model fit. Figure 5.10, Figure 5.11, and Figure 5.12 show how many times the AIC, BIC, and adjusted BIC, respectively, selected each model reduction as the preferred model for a given data set.
Figure 5.10 shows a strong preference for the saturated model when using the AIC to pick the preferred model. This indicates that the additional parameters in the saturated model provide enough improvement to model fit to justify the additional complexity. However, structural reduction was also selected fairly often, especially in the true Q-matrix conditions. This suggests that the full structural model was often overly complex, and a reduced version provided adequate fit.

The BIC shows very different results (Figure 5.11). When selecting a model based on the BIC, the preferred method was most commonly structural-measurement, followed by measurement-
structural reduction. This may be unsurprising, as the penalty for additional parameters is greater in the BIC than the AIC (Wit, van den Heuvel, & Romeijn, 2012). Therefore, the BIC is more likely to prefer models with fewer parameters. However, when the Q-matrix is correctly specified, the preference for the saturated model increases as the sample size increases. This indicates that model reduction may be more important when sample sizes are smaller, and it is more difficult to get good parameter estimates.

![Figure 5.11: Number of selections as best fitting model as measured by the BIC when reducing using p-values](image)

Finally, results when using the adjusted BIC show a middle ground between the results when
using the AIC or BIC (Figure 5.12). There is still a strong preference for the saturated model, however, there is also a stronger preference for the structural-measurement and measurement-structural reductions than what was seen when using the AIC.

![Figure 5.12: Number of selections as best fitting model as measured by the adjusted BIC when reducing using p-values](image)

5.1.5 Description of reduced parameters

Another outcome of interest beyond the recovery of the generated parameters is how often each reduction process resulted in the correct parameters being included in the final model. For example,
if an item measured only one attribute, but the Q-matrix specified two attributes, were the additional main effect and interaction term correctly removed? Conversely, if an item did in fact measure two attributes, were both main effects and the interaction term correctly retained? Figure 5.13 shows the proportion of time the measurement model was correctly reduced, and Figure 5.14 shows the proportion of correct reductions for the structural model.

Figure 5.13: Proportion of correct measurement model reductions when reducing with p-values
Figure 5.14: Proportion of correct structural model reductions when reducing with p-values

Figure 5.13 shows that the only cases where the correct measurement model was obtained was when it was correctly specified, and no parameters were removed (i.e., saturated model or only structural reduction). These results should be interpreted with caution. A “correct” measurement model is defined as all parameters in the final model exactly matching the parameters used to generate the data. This does not account, however, for parameters that may have contributed to the data generation, but not in a significant way. For example, the main effects were drawn from a uniform distribution (0.00, 5.00]. Thus, it is entirely plausible, and expected, for main effects with a value close to zero to be drawn. Thus, these parameters would technically be part of the
true measurement model, but may not be statistically significant, and even one of these parameters would result in the determination of an incorrect measurement model.

A similar pattern is seen in Figure 5.14, with very low rates of correct reduction, except when no reduction of the structural model was performed (i.e., saturated model or only measurement reduction). Because the data was always generated with a fully saturated log-linear structural model, and reduction of the structural model results in a determination of an incorrect reduction. However, given that the structural parameters were all drawn from a $\mathcal{N}(0, 2)$ distribution the same caveats of significance apply that applied to the measurement model parameters.

To verify that the parameters being reduced are indeed these parameters that are very small, and thus may not be significant, the distributions of estimates and standard errors of the reduced parameters can be examined. Figure 5.15 shows that the majority of reduced measurement model parameters are small in value, with small standard errors. More parameters were removed from the true Q-matrix conditions than the over specified Q-matrix conditions, however this is an artifact of the over specified conditions converging far less often.

Figure 5.16 shows a similar pattern for the structural parameters, with the majority of reduced parameters having an estimate close to zero. Additionally, the most of the reduction occurred out of the true Q-matrix conditions, as the over specification conditions often failed to converge. However, unlike the measurement model parameters, there is more variability in the estimates and standard errors of the parameters that were reduced.
Figure 5.15: Distributions of estimates and standard errors for reduced measurement model parameters when reducing with p-values
Figure 5.16: Distributions of estimates and standard errors for reduced structural model parameters when reducing with p-values

5.2 Reduction by heuristic

When the saturated model failed to converge, high level interaction parameters (i.e., three- and four-way interactions) were selected for reduction from the measurement and structural models. Following this initial heuristic reduction, if the model converged, the reduction processes continued using p-values (Figure 4.1). Whereas section 5.1 described the performance of model that were reduced using p-values at all steps of model reduction, this section describes the second scenario,
where models were reduced with the heuristic first, and then p-values.

5.2.1 Convergence

Figure 5.17 shows the convergence rates for these reduction processes when the saturated model failed to converge. Recall that measurement-structural and structural measurement reduction only occurred if the measurement and structural reductions using the heuristic, respectively, converged. When the Q-matrix is correctly specified, reduction of the structural model led to convergence the majority of the time. Conversely, the removal of three- and four-way interactions in the measurement model never led to convergence, unless the structural was being reduced simultaneously. The measurement model was able to be reduced by p-values, but only after higher order interactions had been reduced from the structural model.
In contrast to the truly specified Q-matrix conditions, models rarely converged when the Q-matrix was over specified, regardless of reduction process. This was especially true as the complexity and amount of over specification increased. However, across these conditions, simultaneous reductions and measurement model reduction had the most success in getting the model to converge.

Figure 5.17: Convergence rates when reducing using a heuristic
5.2.2 Parameter recovery

As with the reduction process that relied entirely on p-values, the bias and mean square error of the parameter estimates can be examined for models that used the heuristic as the first reduction step. Figure 5.18 and Figure 5.19 show the total bias and total mean square error, respectively, across all measurement model parameters when reducing using p-values. As with the Because of some outlying data sets, biases with an absolute value greater than 100 and mean square errors with a value greater than 100 have been excluded from the figures. Figures showing all biases and mean square errors, separated by the type of parameter can be seen in Appendix B.

![Graph showing bias in measurement model main effect estimates](image)

Figure 5.18: Bias in measurement model main effect estimates when reducing using a heuristic
Figure 5.19: Mean square error in measurement model main effect estimates when reducing using a heuristic.

Figure 5.18 shows that across all conditions, as with p-value based reduction, there is relatively little bias in the measurement model parameters, especially when the Q-matrix is correctly specified. The large negative biases seen in the structural-measurement reduction condition for the three attribute and 10 percent over specified Q-matrix are due to a couple of outlying data, as can be seen in Appendix B. Also similar to the reduction processes based entirely on p-values, there is relatively large mean square error values across conditions. As expected, and as with the p-value reduction, this decreases as the sample size increases.
Figure 5.20 and Figure 5.21 show the bias and mean square error of the structural parameters respectively. Overall, there is very little bias and mean squared error in the structural parameters. The instances where larger bias and mean squared error are indicated (e.g., 10 and 20 percent over specified Q-matrix) are instances where very few of the model actually converged. Thus, these values are based on only a few replications. Thus, these results suggest that, as with p-value only reduction, the structural parameters are usually well estimated regardless of model reduction method.

Figure 5.20: Bias in structural model parameter estimates when reducing using a heuristic
Figure 5.21: Mean square error in structural model parameter estimates when reducing using a heuristic

### 5.2.3 Mastery classification

Just as with reduction based on p-values, it is important to evaluate mastery classification (at the attribute and profile level) when the initial reduction step is based on a heuristic. Figure 5.22 and Figure 5.23 show the attribute level agreement as measured by the average correct classification rate and average Cohen's $\kappa$, respectively.
Figure 5.22: Average correct classification rate of attribute mastery when reducing using a heuristic
Figure 5.23: Average Cohen’s κ of attribute mastery when reducing using a heuristic

Across all conditions, both the correct classification rate and Cohen’s κ show high rates of agreement between true and estimated attribute classifications, just as was observed when using p-values for all reductions.

The overall profile classification shows also shows pattern similar to the p-value only reduction (Figure 5.24 and Figure 5.25). The overall profile agreement is consistently lower than the mastery classification at the attribute but, but the profile agreement is generally consistent across all model reduction processes.
Figure 5.24: Average correct classification rate of profile assignment when reducing using a heuristic.
Figure 5.25: Average Cohen’s $\kappa$ of profile assignment when reducing using a heuristic

5.2.4 Model fit

When the saturated model fails to converge and the model reduction process uses a heuristic to remove parameters, it is still important to assess model fit. Figure 5.26 shows the results when using the AIC to select the preferred model. When the Q-matrix is correctly specified, there is a strong preference for structural reduction over structural-measurement reduction. In contrast, when the Q-matrix is over specified, there is a stronger preference for simultaneous reduction of structural and measurement models. This preference is stronger as the amount of over specification
increases.

Figure 5.26: Number of selections as best fitting model as measured by the AIC when reducing using a heuristic

The BIC results in Figure 5.27 are similar to those seen when reduction depends only on p-values. There is a much stronger preference for structural-measurement and measurement-structural reduction than when using AIC. However, the difference is not as pronounced as with p-value only reduction, as simultaneous reduction is still preferred overall. Further, just as with the AIC, the preference for measurement reduction increases as the amount of over specification increases.
The model selection when using the adjusted BIC is also reflective of the results found when using only p-values (Figure 5.28). That is, the adjusted BIC serves as a middle ground between the AIC and the BIC. There is a stronger preference for the structural-measurement and measurement-structural reduction processes than when using the AIC, but not as strong when using the BIC. Additionally, there is a stronger preference for measurement model reduction as the amount of over specification in the Q-matrix increases.
Figure 5.28: Number of selections as best fitting model as measured by the adjusted BIC when reducing using a heuristic.

### 5.2.5 Description of reduced parameters

Finally, the performance of the different processes in terms of correctly specifying the parameters to be included can be examined. Figure 5.29 shows the proportion of time the measurement model was correctly reduced, and Figure 5.30 shows the proportion of correct reductions for the structural model.
Figure 5.29: Proportion of correct measurement model reductions when reducing with a heuristic.
Figure 5.30: Proportion of correct structural model reductions when reducing with a heuristic

Figure 5.29 shows that the only cases where the correct measurement model was obtained was when it was correctly specified, and no parameters were removed using p-values (i.e., simultaneous, measurement, structural, or measurement-structural reduction). However, this should be interpreted with caution. When the Q-matrix is correctly specified, there are no three- and four-way interactions, thus, the heuristic has no parameters to remove. When the Q-matrix is over-specified, removing three- and four-way interactions does nothing to fix the additional main effects that have been added to the model.

Additionally, Figure 5.30, shows that the structural model most usually only correct when
only the measurement model was reduced. This is because the data was always generated with a fully saturated structural model. Thus, any removal of parameters (including three- and four-way interactions) results in an incorrect final model.
Chapter 6

Conclusions

The present study utilizes Monte Carlo simulation to evaluate the performance of various model reduction processes for diagnostic classification models. There are two components of DCMs that can be reduced during the estimation process: the measurement model and the structural model. The measurement model governs how items related to the attributes, and the structural model controls the base rate probabilities of inclusion in each class. In this study, the relative strengths and weaknesses of reducing each model, and in which order was examined. Specifically, convergence rates, the bias and mean square error of the parameter estimates, and the attribute and profile mastery classifications were used to evaluate the processes.

Parameters were chosen for reduction based on their p-value. If after estimation of the prior model in the reduction process the parameter was non-significant (i.e., p-value greater than .05), it was removed from the model in the next iteration. In addition, parameters could also be reduced using a heuristic if the saturated model failed to converge. In practice, practitioners are unlikely to stop if the saturated model fails to converge, and instead remove parameters in an effort to achieve a converged model (e.g., Bradshaw et al., 2014). Thus, if the saturated model failed to converge, three- and four-way interaction terms were removed from the measurement and/or saturated models for the reduction. Following the initial heuristic decision, subsequent reductions were performed based on p-values of converged models. These two processes (p-value only reduction and heuristic and p-value reduction) were analyzed separately to evaluate possible differences in reduction preference.

Overall, the results suggest that all model reduction processes provide reasonably unbiased estimates of measurement model and structural model parameters, and mean square error decreased
as the sample size increased. Additionally, all reduction processes showed high levels of agreement between the true and estimated attribute mastery of the respondents. This was especially at the attribute level as compared to the profile level, but showed a consistent pattern across both p-value based and heuristic based reduction.

However, a key difference found in this study was the convergence rates of different model reduction processes, and how those were affected by the initial convergence of the saturated model. When the saturated model converged, reducing the measurement model first was most likely to lead to a converged model, especially if the Q-matrix was over specified. In contrast, when the saturated model failed to converge, reducing the structural model was far more likely to lead to model convergence. This was most pronounced in the correctly specified Q-matrix conditions. Regardless of whether or not the saturated model converged, model reductions were unlikely to result in a converged solution when the Q-matrix was over specified. The overall low rate over convergence for the over specified Q-matrices, across all conditions, indicates that the model is highly unlikely to converge if the Q-matrix has even low rates of over specification.

Additionally, the model fit results suggest other differences. When the saturated model successfully converged, that was most often the preferred model. Obviously when the saturated model fails to converge, this is not an option. In this scenario, structural reduction was the preferred method of model reduction. It is possible that this finding for the heuristic reduction is an artifact of the structural reduction converging the most frequently. However, structural reduction was also the second most preferred reduction process when reducing with p-values. Together, this suggests that reduction of the structural model is less likely to negatively impact model fit on average.

Taken in totality, the results of this study suggest that the path of model reduction should be determined by the convergence of the saturated model. Should the saturated model converge, then the measurement model should be reduced first in order to create a more parsimonious model, while still maintaining a converged solution. On the other hand, if the saturated model fails to converge, reducing the structural model is most likely to provide a converged solution to evaluate. If this too fails to converge, then the most likely scenario is that the Q-matrix has been misspecified,
and should be revised with input from content area experts, just as in the process for developing the Q-matrix (Bradshaw, 2017).

### 6.1 Limitations and future directions

There are several limitations of this study that deserve additional investigation in future work. First, reduction of converged models was based on p-values. The reliance on p-values is problematic for several reasons. For example, p-values do not give direct inferences about the parameter values, and they are unable to provide information about the actual size of the effect (see Wasserstein & Lazar, 2016 for a more complete summary). However, because of the .05 cutoff used to identify non-significant parameters for reduction, there is likely a non-negligible amount of Type I error. Instead, it would likely be better to use a Bayesian estimation, where credible intervals could be formed around the parameter estimates, and reduced based on the proportion of the posterior distribution that is below a predefined cutoff of practical significance. Improvements in available software for estimating DCMs is likely necessary in order for this to be a viable path forward.

Secondly, the random generation of structural parameters resulted in respondents being unevenly placed into classes. Although this may be more reflective of reality, where some classes are less likely than others (for example Figure 3.2), it also likely contributed to convergence problems. Thus, future research may benefit from having a fixed structural model, or least utilizing some mechanism of ensuring that respondents are present in all classes. As an example, it would be relatively straight forward when generating the structural parameters to keep regenerating parameters until a set is generated that results in all classes having a base rate probability above a given threshold.

Related to the issues of under represented classes, part of the model reduction process could involve removing classes that have a low number of respondents assigned to them (e.g., Templin & Bradshaw, 2014a). The implementation of this process would present several challenges. For instance, without theoretical support for why a given profile may or should not exist in the population, there is no straight forward method for assessing whether or not a profile should be removed.
from the model. In other words, there is a problem in deciding how large a class must be in order for it to be retained. The base rate probabilities could be used, but then a determination must be made about what percentage constitutes a non-significant proportion of the population. Despite these complications, this remains an important area of future research.

Finally, it should be noted that the model selection process should not merely be an exercise in finding a path to convergence. The findings of this study are largely driven by the convergence rates of various model reduction processes. Although this is an important outcome measure, a model should not be selected for use due solely to the fact that it was able to converge. Rather, additional and more accurate measures of model fit are necessary to support the use of a model. Mplus provides $\chi^2$ statistics for univariate and bivariate sets of items; however, these are unable to sufficiently assess model fit due to the violations of the asymptotic assumptions of the distributions. Instead, model fit may be better assessed through posterior predictive model checks from a Bayesian estimation (e.g., Gelman & Hill, 2006; Gelman et al., 2014). In this way, the practitioner can evaluate the overall fit of the model to data, not just compare between models that were able to successfully converge.

Despite these limitations, this study provides the first empirical evaluation of model reduction processes in DCMs. As the operational use of these models continues to grow, continued research into the practical applications of DCMs will need to keep pace. This study not only demonstrates a framework for future research into the application of DCMs, but also provides guidance to researchers and practitioners as to how best to proceed with model estimation.
References


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Templin, J., & Bradshaw, L. (2014a). Hierarchical diagnostic classification models: A family of


mar of graphics.


Appendix A

Parameter Recovery with Reduction from P-values

A.1 Individual Bias

Figure A.1: Bias in measurement model intercept estimates when reducing using p-values
Figure A.2: Bias in measurement model main effect estimates when reducing using p-values
Figure A.3: Bias in measurement model 2-way interaction estimates when reducing using p-values
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Figure A.4: Bias in measurement model 3-way interaction estimates when reducing using p-values
Figure A.5: Bias in measurement model 4-way interaction estimates when reducing using p-values
Figure A.6: Bias in structural model estimates when reducing using p-values
A.2 Individual Mean Square Error

Figure A.7: MSE in measurement model intercept estimates when reducing using p-values
Figure A.8: MSE in measurement model main effect estimates when reducing using p-values
Figure A.9: MSE in measurement model 2-way interaction estimates when reducing using p-values
Figure A.10: MSE in measurement model 3-way interaction estimates when reducing using p-values
Figure A.11: MSE in measurement model 4-way interaction estimates when reducing using p-values
Figure A.12: MSE in structural model estimates when reducing using p-values
Appendix B

Parameter Recovery with Reduction from Heuristic

B.1 Individual Bias

![Bias in measurement model intercept estimates when reducing using a heuristic](image)

Figure B.1: Bias in measurement model intercept estimates when reducing using a heuristic.
Figure B.2: Bias in measurement model main effect estimates when reducing using a heuristic
Figure B.3: Bias in measurement model 2-way interaction estimates when reducing using a heuristic
Figure B.4: Bias in measurement model 3-way interaction estimates when reducing using a heuristic
Figure B.5: Bias in measurement model 4-way interaction estimates when reducing using a heuristic
Figure B.6: Bias in structural model estimates when reducing using a heuristic
### B.2 Individual Mean Square Error

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Figure B.7: MSE in measurement model intercept estimates when reducing using a heuristic
Figure B.8: MSE in measurement model main effect estimates when reducing using a heuristic
Figure B.9: MSE in measurement model 2-way interaction estimates when reducing using a heuristic.
Figure B.10: MSE in measurement model 3-way interaction estimates when reducing using a heuristic
Figure B.11: MSE in measurement model 4-way interaction estimates when reducing using a heuristic
Figure B.12: MSE in structural model estimates when reducing using a heuristic