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Inter-rater Reliability of the McKenzie System of Mechanical Diagnosis and Therapy in the Examination of the Knee

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A thesis submitted in partial fulfillment of the requirements for the degree in Master of Science

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INTER-RATER RELIABILITY OF THE MCKENZIE SYSTEM OF MECHANICAL DIAGNOSIS AND THERAPY IN THE EXAMINATION OF THE KNEE

(Thesis format: Monograph)

by

Sean Willis

Graduate Program in Health and Rehabilitation Sciences

A thesis submitted in partial fulfillment of the requirements for the degree of Master of Science Health and Rehabilitation Sciences (Physical Therapy)

The School of Graduate and Postdoctoral Studies
The University of Western Ontario
London, Ontario, Canada

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Abstract

Objective: The purpose of this thesis was to investigate the inter-rater reliability of the McKenzie System of Mechanical Diagnosis and Therapy (MDT) when classifying patients with musculoskeletal knee pain using clinical vignettes. Methods: This study was divided into two phases. First, ten clinicians experienced in the use of MDT were randomly recruited to write a total of 60 clinical vignettes based upon the initial assessment of past patients with knee pain. Second, six different MDT raters were recruited to rate 53 selected vignettes and reliability was determined using Fleiss Kappa. Results: There was “substantial agreement” among six MDT raters classifying the clinical vignettes into one of four categories (kappa=0.72). There was no statistically significant difference between therapists with different levels of training. Significance: These findings indicate that the McKenzie System of MDT is a reliable method of classifying patients presenting with musculoskeletal knee pain when using clinical vignettes.

Keywords

McKenzie system, knee, Mechanical Diagnosis and Therapy, reliability, vignette
Co-Authorship Statement

Each of the four chapters in this thesis is based upon research designed and organized by the author. Members of the advisory committee provided regular guidance and feedback during this process. All chapters were authored primarily by Sean Willis with editing provided by the advisory committee.
Acknowledgments

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List of Abbreviations

ACL – anterior cruciate ligament

CI – confidence interval

LR – likelihood ratio

MDT – McKenzie System of Mechanical Diagnosis and Therapy

MRI – magnetic resonance imaging

NBA – National Basketball Association

OA – osteoarthritis

OSTs – orthopaedic special tests

PFPS – Patellofemoral Pain Syndrome

QUADAS – Quality Assessment of Diagnostic Accuracy Studies

SD – standard deviation

SE – standard error
Chapter 1

1 Overview of Problem

In the US, the prevalence of knee pain has increased by 65% over the last 20 years (Nguyen et al., 2011). Over 4 billion dollars are spent annually on arthroscopic knee surgery alone (Gage, McIlvain, Collins, Fields & Comstock, 2012) despite evidence suggesting arthroscopic surgery does not result in superior patient outcomes (Kirkley et al, 2008; Sihovnen et al, 2013; Thorlund, Juhl, Roos & Lohmander, 2015). Osteoarthritis (OA) related knee pain has been identified as a possible trigger for physical and functional decline for older adults (Jinks, Jordan & Croft, 2007). For those suffering from OA related hip and knee pain the cost of time lost from employment and leisure as well as their unpaid caregivers is often underestimated in the contribution to the overall burden of OA (Gupta, Hawker, Laporte, Croxford & Coyte, 2005). Within the current environment of fiscal responsibility in healthcare, it is vital that the overall costs of knee pain are recognized so that interventions that reduce the physical and financial burden are identified and funded to maximize patient outcomes. For this to occur, clinicians must possess the skills or use methods of assessment that have the clinical utility to identify the most appropriate, cost effective intervention from which the patient will benefit.

Therefore, it is essential that an orthopaedic evaluation of the knee is valid and reliable, is guided by clear diagnostic criteria and provides the clinician with prognostic value.

An established body of evidence highlights limitations with the diagnostic validity of orthopaedic special tests (OSTs) used in the clinical examination of the knee (Cook, Mabry, Reiman & Hegedus, 2012; Geraets et al., 2015; Hegedus, Cook, Hasselblad, Goode & McRory, 2007; Lange et al., 2014; Leblanc et al., 2015; Peeler, Leiter, & MacDonald, 2010). The reported psychometric properties and hence the diagnostic accuracy of many of the commonly used OSTs are influenced by a number of factors including but not limited to rater experience, varied interpretation of the result findings, lack of a standardized approach to performance of the test and study design-related bias (Cook et al., 2012; Hegedus et al., 2007; Geraets et al., 2015; Lange et al., 2014; Leblanc et al., 2015; Peeler et al., 2010). Moreover, research around medical imaging such as
magnetic resonance imaging (MRI) have investigated the limitation of findings through the presence of asymptomatic pathology and abnormalities (Beattie et al., 2008; Boks, Vroegindeweij, Koes, Hunink & Bierma-Zeinstra, 2006; Kaplan, Schurhoff, Selesnick, Thorpe & Uribe, 2005; LaPrade, Burnett, Veenstra & Hodgman, 1994). With the relationship between pain and radiographic pathology not fully understood and limitations in reported diagnostic accuracy of OSTs, it has been suggested that specifically defined criteria used by non-pathoanatomical classification systems may offer better utility and should be considered as an alternative to the current model (Rosedale et al., 2014).

One system that has not been thoroughly tested for use with musculoskeletal pain in the extremity is the McKenzie System of Mechanical Diagnosis and Therapy (MDT). The MDT system of classification uses a non-pathoanatomically specific approach to classify patients based on their response to repeated end range loading strategies. Although demonstrating good inter-rater reliability in the assessment of musculoskeletal spinal pain (Clare, Adams & Maher, 2005; Kilpikoski, Airaksinen, Kankaanpaa, Leminen, Videman & Alen, 2002; Razmjou, Kramer & Yamada, 2000), shoulder pain (Heider Abady, Rosedale, Overend, Chesworth & Rotondi, 2014) and the extremities (Kelly, May, & Ross, 2008; May & Ross, 2009), MDT has not been evaluated on its use in the assessment of musculoskeletal knee pain.

Clinical vignette based methodologies are often used in the evaluation of decision making and clinical judgment of health professionals (Evans et al., 2015). Although often criticized because they do not reflect actual practice which may influence results and conclusions of studies, well designed vignette studies can be practical, offer flexibility, avoid ethical and observational issues and be generalizable to real world settings (Evans et al., 2015; Peabody, Luck, Glassman, Dresselhaus & Lee, 2000; Rutten, Harting, Rutten, Bekkering & Kremers, 2006). Clinical vignettes are an inexpensive option to control multiple variables, collect information simultaneously from multiple sources, and isolate clinical decision making. Thus they can provide an initial step in the investigation of the reliability of MDT for knee conditions.
1.1 Purpose

Considering the classification system of MDT has not been rigorously tested in the assessment of musculoskeletal knee pain and the use of clinical vignette based methodologies are a valid approach to examine clinical decision making, the purpose of this study is to determine the inter-rater reliability of MDT in the examination of the knee and what influence the level of MDT training may have on reliability.

1.2 Structure of Thesis

This document is presented in the “monograph” format described by the Western University Faculty of Graduate Studies.

In Chapter 2, a review of the literature is performed examining medical imaging and the presence of asymptomatic pathology, OSTs in the examination of the knee, the validity and reliability of MDT and the use of clinical vignettes in medical research.

Chapter 3 outlines the methods and presents results of the study. Chapter 4 discusses study findings, implications for health care professionals and limitations of the study. Recommendations for future research in this area are outlined.
Chapter 2

2 Literature Review

This chapter reviews the key findings and conclusions from a literature review in the areas of medical imaging and asymptomatic pathology, orthopaedic special tests (OSTs) in the examination of the knee, the McKenzie system of Mechanical Diagnosis and Therapy and the use of clinical vignettes in the research of clinical decision making. Gaps in the current research are also identified.

2.1 Medical Imaging and Asymptomatic Pathology

The diagnostic accuracy of an orthopaedic test is dependent on its ability to rule in or rule out pathology. The clinical utility of that test may be partially determined by the ability of that test to discriminate between symptomatic and asymptomatic pathology or abnormalities. Studying athletes and active individuals, a number of articles have highlighted the presence of previously undiagnosed anatomical abnormalities with medical imaging in pain-free individuals (Beattie et al., 2008; Boks et al., 2006; Kaplan et al., 2005; LaPrade et al., 1994).

Kaplan et al. (2005) reviewed the knee MRI findings of 20 National Basketball Association (NBA) players that met the inclusion criteria of no history of knee pain or surgery and had negative tests on physical examination for the presence of knee abnormalities such as meniscal and ligamentous disruptions and patella-femoral joint pain. The findings of the study, looking at players ranging from 21 to 36 years old, found that 47.5% of the evaluated knees had articular cartilage lesions and 20% of knees had meniscal tears. In their conclusion, the authors noted the influence of diagnostic imaging on clinical decisions and cautioned that findings do not indicate symptoms or functional level.

Investigating 100 patients with suspected meniscal tear, Zanetti, Pfirrmann, Schmid, Romero, Seifert and Hodler (2003) found 57 patients on MRI had a meniscal lesion on the symptomatic knee and of those, 36 had a meniscal lesion on the asymptomatic side
(63%). In a similar study, Boks et al. (2006) examined the MRI results of 134 patients with knee pain and found that of the 45 patients with a meniscal tear on the symptomatic side, 19 (42%) had one on the asymptomatic side. These values are substantially higher than those previously reported by LaPrade et al. (1994) who concluded that emphasis is needed on the importance of matching MRI findings with the history and physical examination after finding a prevalence rate of 5.6% for asymptomatic meniscal tear in 54 men and women with no previous history of knee pain or trauma. What should be noted is that Kaplan et al. (2005), LaPrade et al. (1994) and Zanetti et al. (2002), screened subjects for knee pathology prior to their participation.

Using a sample comprised of men and women of an average age of 41.5 years of age, Beattie et al. (2008), recruited subjects with no previous history or diagnosis of knee pathology to undergo a MRI and X-ray on their non-dominant knee. Although the prevalence of cartilage lesions was relatively low at 11%, all but one participant exhibited a meniscal abnormality in at least one region of the knee with more than 60% of participants having an abnormality in at least three of four regions.

In summary, it has been suggested that MRI findings should be interpreted with caution as findings do not indicate symptoms or functional level (Kaplan et al., 2005). With evidence demonstrating the presence of asymptomatic pathology in the knee, it is reasonable to question not only the diagnostic utility of detecting pathology but also whether one can be certain an implicated structure is the cause of an individual’s symptoms. This can be of particular consequence when patients present with a history of pain and MRI identified pathology to which a decision on care must be made, often having to decide whether or not surgery is indicated.

### 2.2 Orthopaedic Special Tests (OSTs) in the Examination of the Knee

A change in practice has gradually occurred over the last several decades as clinicians performing an orthopaedic assessment have become over reliant on the results of OSTs and medical imaging (Cook, 2010). The psychometric properties and hence, the diagnostic accuracy of these OSTs is often influenced by a number of factors including
but not limited to rater experience, varied interpretation of the result findings and lack of a standardized approach to performance of the test (Cook et al., 2012; Hegedus et al., 2007; Peeler et al., 2010). Several threats to diagnostic validity such as selection bias, verification bias and the study sample have been identified and can inflate estimates of diagnostic accuracy (Lijmer et al., 1999). As a result, the usefulness of many of these tests has been questioned (Cook et al., 2012; Geraets et al., 2015; Goossens et al., 2015; Hegedus et al., 2007).

Investigating the diagnostic accuracy of three common ACL tests, Peeler et al. (2010) found only moderate levels of inter-rater agreement for the anterior drawer (0.57), the Lachman (0.45), and the pivot shift (0.53). The Lachman demonstrated a sensitivity of 83% with orthopaedic surgeons but varied greatly within clinician groups, family physicians and therapists, ranging from 15% to 87%. Peeler et al. concluded that variables such as level of experience and degree of training or specialization may impact the accuracy of testing. Geraets et al. (2015) had similar findings comparing an orthopaedic surgeon and primary care physician and the diagnostic value of the subjective and objective exams. They concluded that the objective exam, while improving an orthopaedic surgeons’ positive predictive value of an ACL tear from 0.65 to 0.94, offered no value to the assessment for the primary care physician, dropping positive predictive value from 0.69 to 0.62.

Leblanc et al. (2015) have suggested that the clinical setting and the degree of tear will impact the diagnostic accuracy of tests for anterior knee instability. In their systematic review, they found that the sensitivity of the Lachman and pivot shift tests were lower when patients were awake versus under anesthetic and in the presence of a partial versus a complete tear. They also found insufficient data to calculate a pooled specificity and as a result, were unable to give a clear recommendation of the diagnostic accuracy of the physical examination in ACL deficient knees. Similarly, Lange et al. (2015) were unable to perform a meta-analysis during their systematic review of the physical tests for ACL rupture as a result of heterogeneity of the sample populations, the reliability measures used and the poor methodological quality of the studies reviewed.
Of the 18 studies qualifying to be included in a meta-analysis of the physical tests for meniscal tears, Hegedus et al. (2007) found three tests to be studied most: McMurray’s, Apley’s and joint line tenderness. Of those three tests, McMurray’s demonstrated the highest sensitivity of 70% but also the lowest specificity at 71%. Joint line tenderness had the highest specificity with 77% but also the lowest sensitivity at 63%. Hegedus et al. concluded that no single test is able to accurately diagnosis a meniscal tear and recommended that the performance and interpretation of the tests be standardized.

More recently, Goossens et al. (2015) reported the Thessaly meniscal test with a sensitivity of 64% and a specificity of 53%. When combined with the McMurray’s test, the sensitivity dropped to 53%. As a result, the authors concluded that either test in isolation or combined, does not appear useful in the detecting of meniscal tears. Further to this, it was recommended that research should focus on the development of a better diagnostic model of examination. Of interest, Campbell et al. (2014) investigated the correlation between location of preoperative knee pain and arthroscopic knee findings. The authors found that no significant correlation (p=0.98) existed between pain location and pathology and concluded that because of the varied nature of pain their results dispute the widely held beliefs that the location of pain is related to underlying pathology.

Cook et al. (2012) conducted a systematic review for clinical tests for screening and diagnosing patellofemoral pain syndrome (PFPS) using the Quality Assessment of Diagnostic Accuracy Studies (QUADAS) scoring for methodological quality. Of the 704 articles identified, 9 met selection criteria, presenting with 22 clinical tests for review. None of the 22 tests reviewed demonstrated a positive likelihood ratio (+LR) greater than 5.0 and a negative likelihood ratio (-LR) less than 0.20, meaning an inability to rule in or out (PFPS). Of those tests that had a stand-alone +LR greater than 5.0, those studies had the lowest methodological quality and/or used normals as the control group introducing quality bias and affecting diagnostic accuracy.

In summary, while widely accepted, the use of OSTs in the examination of the knee demonstrates limited ability to establish a clear relationship between clinical testing and symptomatic pathology. A potential explanation for this limitation may be the over-
reliance on identifying an anatomical structure or structures that are the cause of pain. For the ideal management of musculoskeletal problems, clinicians require the use of accurate tests and validated diagnostic criteria. Issues with the current approach highlight the need to explore other systems of clinical examination.

2.3 McKenzie System of Mechanical Diagnosis and Therapy

The McKenzie System of Mechanical Diagnosis and Therapy (MDT) is a non-pathoanatomically specific classification system that was originally developed for use in spinal conditions. The assessment screens out potential red flag issues such as fractures, neurological or vascular issues to determine if a patient’s symptoms are mechanical in nature. MDT involves a detailed history and an examination in which baseline symptoms, both with function and at rest, are established and then re-evaluated following the patient performing repeated end range loading movements to the affected area. A key characteristic of the system that has shown potential as a prognostic indicator of musculoskeletal pain is directional preference (May & Aina, 2012). Directional preference is defined as the rapid improvement of a patient’s symptoms with positioning or movement in one specific direction while commonly worsening with positioning or movement in the opposite direction (McKenzie and May, 2003). Based on the patients’ response to the assessment and potential change in baseline symptoms, the clinician is able to formulate a provisional classification and provide directed treatment.

The system is based not on determining an anatomical diagnosis but rather classification into one of four categories, the first three being specific mechanical syndromes: Derangement, Dysfunction, Postural or OTHER. The mechanical syndromes were originally developed based on particular patterns of symptoms and responses to movement in the spine that were seen by the founder of MDT, Robin McKenzie. More recently, the MDT system has been used with increased frequency by trained clinicians in the evaluation of joints in the extremities.

The Derangement syndrome is the most common of the three mechanical syndromes and is varied in its clinical presentation; however, the key characteristic is the presence of a
directional preference with loading strategies (McKenzie and May, 2003). An example of this would be a worsening of a patient’s symptoms with movements into flexion but an improvement or abolishment of symptoms with movements into extension. The clinical presentation for Dysfunction syndrome is intermittent pain that is consistently reproduced at the end-range of a restricted movement but will not persist once mechanical loading strategies have ceased (McKenzie and May, 2003). In the extremities, the Dysfunction syndrome can be disseminated further as Articular and Contractile Dysfunction. Contractile Dysfunction is characterized by pain brought on by active and resisted movements, where passive range of motion is generally preserved (McKenzie and May, 2000). Articular Dysfunction is distinguished from contractile through the loss of active and passive range of motion with pain being produced at the end of available range and absent during resisted testing (McKenzie and May, 2000). Postural syndrome is distinguished by local, intermittent pain without movement loss that is brought on by sustained postures and abolished with posture correction (McKenzie and May, 2003). An annual review of the MDT educational program is conducted by the International Education Committee of the McKenzie Institute International. Any revisions or changes to the definitions or criteria are brought about by updates to published research literature, through feedback from MDT Faculty and of the evaluation of the system from the Committee members. The most current summary of the classifications is provided in Table 1 and is presently in use in the MDT education manuals. The OTHER category is made up of 10 diagnostic subgroups which together complete the full MDT classification system where each subgroup has its own definition and diagnostic criteria (Table 2). Table 2 has been modified from the original publication by May and Rosedale (2012) to reflect the most recent revisions made by the International Education Committee.

The MDT system of education has two levels of clinical competence, Credentialed and Diploma. Credentialed clinicians have completed four post-graduate courses and successful passed a standardized written and practical examination. Having attained Credentialed status, clinicians can then go on to acquire Diploma which consists of one University semester theoretical component and 360 hours of clinical practice mentorship. Once completed, the clinician must then pass an oral examination to be awarded Diploma status (http://www.mckenzieinstitute.org/).
In the literature, a spinal assessment using MDT has been shown to have good inter-rater reliability (Clare et al., 2005; Kilpikoski et al. 2002; Razmjou et al., 2000). Using two MDT trained therapists, one Credentialed and one Diploma, Razmajou et al. (2000) investigated the interrater reliability of the MDT system during the assessment of 45 patients presenting with mechanical low back pain. They found the overall reliability between raters on mechanical syndrome classification to be substantial (kappa=0.70) with the Derangement classification to have the highest reliability of kappa=0.96. The agreement of syndrome classification was 93% between raters for all responses. These results are similar to those of Kilpikoski et al. (2002) that found an overall reliability between two MDT trained Diploma raters on the assessment of 39 patients to be moderate (kappa=0.6) with 95% agreement on syndrome classification. The majority of participants (90%) were classified into the Derangement syndrome.

Clare et al. (2005) examined the reliability of the MDT system in the classification of patients with lumbar and cervical pain between 14 raters, seven Credentialed and seven Diploma. The overall reliability for the classification of patients was substantial (kappa=0.84) with kappa=1.0 for lumbar patients and kappa=0.63 for cervical patients. Overall agreement amongst raters was high with 96% for the total patient pool. The majority of patients (66%) were again classified as Derangement syndrome.

More recently, Werneke et al. (2014) investigated the relationship between pre-Credentialed level of training and therapist agreement in the McKenzie lumbar classification. Forty-seven raters of various levels of pre-Credentialed MDT training assessed over 1600 patients and found an overall range of kappa=0.37 to 0.44 for classification into one of the mechanical syndromes despite an observed agreement of 86 to 91%. It has been suggested this paradox results from the sensitivity of the kappa statistic when the prevalence of a rating is either very high or very low and that the interpretation and reporting of the kappa statistic alone may result in conclusions that may be misleading (Bryt, Bishop & Carlin, 1993; Cicchetti & Feinstein, 1990; de Vet, Mokkink, Terwee, Hoekstra and Knol, 2013; Feinstein & Cicchetti, 1990). Indeed, for patients classified into one of the four mechanical syndrome classifications, Werneke et al. (2014) reported Derangement syndrome among examiners to range from 334 to 512
(81 to 86%) with the remaining three classifications ranging from 0 to 27 (0 to 4.6%). This skewed distribution elevates the probability of agreement due to chance alone and thereby lowers the value of the kappa statistic which represents the proportion of agreement greater than that expected by chance (O’Leary et al., 2014; Werneke et al., 2014).

Previously published studies investigating the value of MDT guided treatment in the extremity had been restricted to case studies documenting patients presenting with shoulder and knee pain (Aina & May, 2004; Littlewood & May, 2007; Lynch & May, 2013) and despite an increase in clinical use, research evaluating the MDT assessment for musculoskeletal extremity problems is limited. In a pilot study, Kelly, May & Ross (2008) examined the reliability of trained MDT clinicians classifying clinical vignettes based on patients with musculoskeletal disorders in the extremity. They found the agreement among three Credentialed raters for 11 vignettes to be kappa=0.70. These results are similar to May & Ross (2009) which investigated the reliability of the MDT assessment form for extremity conditions by using 25 clinical vignettes. They found an overall level of agreement to be 92% with a kappa of 0.83 among 97 Diploma trained therapists. There was little difference in reliability between upper (kappa=0.85) and lower extremity (kappa=0.80) cases.

Surveying Diploma therapists, May and Rosedale (2012) gathered data on the prevalence of mechanical syndromes and treatment strategies in use for the extremities. The most commonly used classifications for patients presenting with musculoskeletal knee pain were: Derangement (42.7%), Articular Dysfunction (3.9%), Contractile Dysfunction (8.7%) and OTHER (44.7%), 20% of which were post-surgery or post-trauma. Of interest, May and Rosedale (2012) found that 85.8% of initial classifications remained stable throughout the treatment episode. More recently, Heider Abady et al. (2014) demonstrated almost perfect reliability (kappa=0.90) between six Diploma raters when using MDT to classify 54 clinical vignettes of patients with musculoskeletal shoulder pain with an overall level of multi-rater agreement to be 96%. Of note, the highest level of agreement in this study was for Spinal, with the category of OTHER having the lowest level of agreement.
In a recent randomized control trial, Rosedale et al. (2014) sought to examine the effectiveness of exercise intervention determined through an MDT on patients diagnosed with end stage knee OA. Not only were patients readily classified as Derangement or not Derangement, but it could be inferred that the large effect size of $d = 0.77$ to $0.87$ for all primary outcomes seen at two weeks by the intervention group is attributable to the classification and exercise matching determined by the MDT assessment. Although the results of this study are encouraging, the reliability of the MDT classification system had not yet been previously studied in the knee.

In summary, although the MDT system has been shown to be reliable for assessment of musculoskeletal pain in the spine and shoulder, no studies have been conducted on the reliability of the MDT system for musculoskeletal knee pain.
<table>
<thead>
<tr>
<th>MDT Classification</th>
<th>Clinical Presentation</th>
</tr>
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<tbody>
<tr>
<td>Derangement</td>
<td>Varied in its clinical presentation; associated with mechanical obstruction of an affected joint; however, the key characteristic is the presence of a directional preference with loading strategies. Directional preference is defined as the rapid improvement of a patient’s symptoms with positioning or movement in one specific direction while commonly worsening with positioning or movement in the opposite direction.</td>
</tr>
<tr>
<td>Dysfunction</td>
<td>Intermittent pain that is consistently reproduced at the end-range of a restricted movement but will not persist once mechanical loading strategies have ceased. In the extremities, the Dysfunction syndrome can be disseminated further as Articular and Contractile Dysfunction.</td>
</tr>
<tr>
<td>Articular Dysfunction</td>
<td>Distinguished from contractile through the loss of active and passive range of motion with pain being produced at the end of available range and absent during resisted testing.</td>
</tr>
<tr>
<td>Contractile Dysfunction</td>
<td>Characterized by pain brought on by active and resisted movements, where passive range of motion is generally preserved.</td>
</tr>
<tr>
<td>OTHER</td>
<td>Category is made up of 10 diagnostic subgroups which together complete the full MDT classification system where each subgroup has its own definition and diagnostic criteria. (Table 2)</td>
</tr>
</tbody>
</table>
Table 2 Subgroups of MDT OTHER classification

<table>
<thead>
<tr>
<th>Serious Pathology (list not exhaustive)</th>
<th>Category</th>
<th>Clinical Findings (Red Flags)</th>
<th>Clinical Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cancer</td>
<td>Age &gt;55, history of cancer, unexplained weight loss, progressive, not relieved by rest</td>
<td>Maybe primary site or metastases</td>
<td></td>
</tr>
<tr>
<td>Fracture</td>
<td>History of significant trauma (If osteoporosis present; minor trauma)</td>
<td></td>
<td>Loss of function. All movements make symptoms worse.</td>
</tr>
<tr>
<td>Infection</td>
<td>Fever, malaise, constant pain, all movements worsen</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Non-Serious Pathology Subgroups for OTHER classification

<table>
<thead>
<tr>
<th>Subgroup</th>
<th>Definition</th>
<th>Criteria</th>
<th>Clinical Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chronic Pain Syndrome</td>
<td>Pain-generating mechanism influenced by psychosocial factors or neurophysiological changes</td>
<td>Persistent widespread pain, aggravation with all activity, disproportionate pain response to mechanical stimuli, inappropriate beliefs and attitudes about pain.</td>
<td>Regional pain syndromes</td>
</tr>
<tr>
<td>Inflammatory</td>
<td>Inflammatory arthropathy</td>
<td>Constant pain, morning stiffness, excessive movements exacerbate symptoms</td>
<td>RA, sero-negative arthritis, some stages of OA</td>
</tr>
<tr>
<td>Subgroup</td>
<td>Definition</td>
<td>Criteria</td>
<td>Clinical Example</td>
</tr>
<tr>
<td>--------------------------</td>
<td>----------------------------------------------------------------------------</td>
<td>----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
<td>------------------------------------</td>
</tr>
<tr>
<td>Mechanically Inconclusive</td>
<td>Unknown musculoskeletal pathology</td>
<td>Derangement, Dysfunction, Postural and subgroups of OTHER excluded. Symptoms affected by positions or movements BUT no recognizable pattern identified OR inconsistent symptomatic and mechanical responses on loading.</td>
<td></td>
</tr>
<tr>
<td>Peripheral Nerve Entrapment</td>
<td>Peripheral nerve entrapment</td>
<td>No spinal symptoms. Local paraesthesia / anaesthesia. May have local muscle weakness.</td>
<td>Carpal tunnel syndrome, myalgia paraesthetica</td>
</tr>
<tr>
<td>Post-surgery</td>
<td>Presentation relates to recent surgery</td>
<td>Recent surgery and still in post-operative protocol period.</td>
<td></td>
</tr>
<tr>
<td>Soft Tissue Disease Process</td>
<td>A fibroblastic or degenerative disease process affecting inert soft tissue with unknown or disputed aetiology</td>
<td>Each disease process has a unique clinical presentation, natural history and response to a variety of interventions.</td>
<td>Frozen shoulder, Dupuytren’s, plantar fascia syndrome</td>
</tr>
<tr>
<td>Subgroup</td>
<td>Definition</td>
<td>Criteria</td>
<td>Clinical Example</td>
</tr>
<tr>
<td>----------------------------------</td>
<td>---------------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------------------------------</td>
<td>----------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Structurally Compromised</td>
<td>Soft tissue and/or bony changes compromising joint integrity</td>
<td>Mechanical symptoms (ROM restricted, clunking, locking, catching).</td>
<td>Late stage OA, dislocation, labral tear, cruciate ligament rupture, irreducible meniscal tear</td>
</tr>
<tr>
<td></td>
<td></td>
<td>May have sensation of instability.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Long history of symptoms or history of trauma.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Irreversible with conservative care.</td>
<td></td>
</tr>
<tr>
<td>Trauma/Recovering Trauma</td>
<td>Recent trauma associated with onset of symptoms</td>
<td>Recent trauma associated with onset of constant symptoms / recent trauma associated with onset of symptoms, now improving and pain intermittent.</td>
<td></td>
</tr>
<tr>
<td>Vascular</td>
<td>Symptoms induced by poor blood supply due to pressure increase in a closed anatomical space.</td>
<td>Below knee symptoms, predominantly in younger athletes.</td>
<td>Compartement syndrome</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Consistently induced by exercise or activity.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>May have pain and /or paraesthesia in field of local cutaneous nerve and local swelling.</td>
<td></td>
</tr>
</tbody>
</table>

2.4 Clinical Vignettes

To assess inter-rater agreement, measurement of a clinicians’ performance must “ultimately rely on measures that are valid, reliable, inexpensive and manageable” (Rutten et al., 2006, p. 492). Two methods presently used in the literature to assess reliability include the use of real patients and clinical vignettes. Each method possesses its own strengths and weaknesses. Recruiting actual patients allows for subtle variability in patient presentation for similar musculoskeletal problems. Actual patients may permit a true expression of the nature of symptoms and responses to testing and potentially allow for better interpretation of the clinical interaction (May & Ross, 2009). Using patients to test inter-rater agreement may improve the realism and depth to the clinical scenario which may improve external validity and generalizability of the study and findings.

However, there are limitations to using real patients. For instance, real patients may make measurement by direct observation difficult to apply, especially in larger samples, can be expensive and time-consuming, and is potentially subject to a Hawthorne effect (Rutten et al., 2006). Use of real patients may result in insufficient case mix (Peabody et al., 2000) which may inadvertently introduce sampling bias, especially as it relates to MDT. The Derangement syndrome is the most common classification in the spine (78%, May, 2006) and in the extremity (37%, May & Rosedale, 2012). Because of the apparent prevalence of the Derangement syndrome, the random recruitment of patients may unintentionally create a homogenous sample that potentially would not include all relevant syndromes within the classification system.

Another option to evaluate inter-rater agreement is through the use of clinical vignettes. Clinical vignettes have a long history of use (Evans et al., 2015) and are defined as written patient case studies based on realistic scenarios where clinicians are given one or more questions asking what they may do if given the actual patient (Veloski, Tai, Evans & Nash, 2005). A number of studies have used vignettes as a primary method of data collection ranging from physical therapy adherence to guidelines (Rutten et al., 2006), best practices of physical and occupational therapist for young patients with cerebral palsy (Saleh et al., 2008), to measuring the quality of physician practice (Peabody et al., 2004). Using clinical vignettes for data collection provide the user the advantages of the
ability to simultaneously collect information from a number of subjects, manipulate multiple variables and create heterogeneous case mixing, avoid ethical issues, and avoid observer effects that can affect observational studies (Gould, 1996).

In a comparison of vignettes, standardized patients and chart abstraction, Peabody et al., (2000) used the three methods to evaluate physician competence and the quality of their practice. The authors concluded that clinical vignettes can be used in an outpatient setting to evaluate quality of care, may offer an inexpensive way to provide adequate case mix and can be a valid and comprehensive means to evaluate processes of care in clinical practice. These findings are consistent with studies by Dresselhaus, Peabody, Luck and Bertenthal (2004) and Veloski et al. (2005), who added that clinical vignettes are an effective way to isolate decision making. In a validation study, Peabody et al. (2004) found clinical vignettes to be a valid tool to provide case-mix variation, and “are particularly useful for comparing quality among and within sites and may be useful for longitudinal evaluations of interventions intended to change clinical practice” (p. 771).

Despite being seen as a valid measurement tool, there appears to be a lack of literature validating framework for the generation or creation of clinical vignettes. When appraising and evaluating articles that use clinical vignettes, Gould (1996) attempted to address this by ensuring certain features were present. Gould (1996) recommended that authors should address internal validity issues by developing vignettes based on existing literature and/or case study review, the scenarios should be tested to remove ambiguity and reviewed by an expert panel that possesses the knowledge and expertise to determine appropriateness of the vignette for the study. Atzmüller & Steiner (2010) proposed that researchers should generate more vignettes than needed and subsequently select those vignettes that would create the best sample with which to test.

Further to this, Veloski et al. (2005) suggested that the clinical scenarios follow the same natural flow of a clinical assessment, that the order of the information be logical and sequenced as though a clinician were performing it on an actual patient. The vignette should be written as such to minimize confusion, remain specific to the goal of testing the hypothetical situation and maintain a level of uncertainty that does preclude the
clinicians’ ability to articulate their interpretation of the case. Well written, realistic vignettes should simulate aspects of real world scenarios, a facet of construct validity, offer enough variability which relates to the study’s internal validity and produce results that are generalizable to real world situations, reflecting external validity (Evans et al., 2015).

2.5 Research Gaps

Although studies have been done to determine the reliability of the MDT system, none have investigated the reliability of the system in the knee. The use of real patients in a reliability study may result in an insufficient case mix which may inadvertently introduce sampling bias, especially as it relates to MDT. With the Derangement syndrome being the most common classification in the spine (78%, May, 2006) and in the extremity (37%, May & Rosedale, 2012), clinical vignettes would allow for the creation of a heterogeneous sample that would include all relevant syndromes within the classification system and thus avoid these issues.
Chapter 3

3 Inter-rater Reliability of the McKenzie System of Mechanical Diagnosis and Therapy in the Examination of the Knee

This chapter reviews the study objectives, design and the methods used to determine the inter-rater reliability of the McKenzie System of Mechanical Diagnosis and Therapy (MDT) in the examination of the knee. The results of the study are also reported in this chapter.

3.1 Study Objectives

The McKenzie System of MDT is a widely used method of classification and management of musculoskeletal problems. Although the McKenzie system has been investigated for its reliability and efficacy in the management of spinal pain, few studies have evaluated the system when applying it to musculoskeletal problems in the extremities, in particular the knee. The objectives of this study were to:

1. To develop 53 clinical vignettes of patients presenting with musculoskeletal knee pain using past patient data from the caseloads of 10 MDT Credentialed or Diploma clinicians which are based on the definitions of four clinical classifications.

2. To test the inter-rater reliability of six MDT-trained experienced clinicians when classifying patients with musculoskeletal knee pain into one of four MDT classifications using written clinical vignettes.

3. To investigate the influence of the level of MDT education on the reliability of classifying patients with musculoskeletal knee pain using written clinical vignettes.

3.2 General Study Design

To achieve these objectives, a two phase study was conducted. To achieve objective one, the first phase consisted of the recruitment of 10 MDT clinicians to develop 53 clinical vignettes representative of the prevalence of musculoskeletal knee pain classified through
MDT. To meet objectives two and three, the second phase required the recruitment of an additional six MDT raters to classify the patients represented in the clinical vignettes and measure the reliability and level of agreement among the MDT raters. Ethical approval for the study was obtained from the Health Science Research Ethics Board at Western University (Appendix A).

3.2.1 Sample Size

Rotondi and Donner (2012) proposed a method of calculating the sample size for studies measuring inter-rater agreement for multiple outcomes and raters. To arrive at an estimated sample size, kappa was set at 0.8 (0.7 lower limit, 0.9 upper limit) based on levels of agreement with previous work evaluating the reliability of MDT in the extremities (Heidar Abady et al., 2014; Kelly, May, & Ross, 2008; May & Ross, 2009). With an alpha of 0.05 for six raters (phase two) and using the prevalence of the four common syndromes of 0.4(Derangement), 0.4(OTHER), 0.1(Contractile Dysfunction), 0.1(Articular Dysfunction) as outlined by May and Rosedale (2012), a value of 53 was determined for the number of clinical vignettes required for phase two. The sample size was estimated using a program developed by Rotondi (2013) for the R Project for Statistical Computing.

3.2.2 Phase 1

3.2.2.1 Participants

For the first phase of the study, ten clinicians experienced in the use of MDT in the extremity were recruited based on previous willingness to participate in research. These clinicians were asked generate 60 clinical vignettes, six vignettes per clinician, classified into one of four classifications. The sample size of ten was chosen to minimize the burden of creating the vignettes on the consenting clinician. To be included, clinicians were Credentialed or Diplomat with the McKenzie institute with more than three years of experience applying MDT to musculoskeletal disorders of the extremity and be registered on the publicly available list of MDT practitioners practicing in the United States or Canada. Clinicians were excluded if they are unable to understand written and spoken English, unable to provide informed consent, or unable to follow the instructions for
generating the clinical vignettes. Correspondence was conducted and informed consent was obtained from each clinician through electronic mail (Appendix B – C). In total, 20 clinicians were approached for recruitment to which 10 consented for participation in this study.

3.2.2.2 Vignette Development

For clinical vignette development, Atzmüller & Steiner (2010), Evans et al. (2015), Gould (1996) and Veloski et al. (2005) have suggested that more vignettes should be generated than will be used, be reviewed by an expert panel to determine appropriateness and to select the best sample for testing, be written based on or relating to a case study or clinical experience and follow a similar structure and natural flow for all vignettes used. For this study, clinicians were asked to generate vignettes based on their past patient assessment files. Clinicians were instructed that each clinical vignette is to be de-identified to only include gender, age range (e.g. 35 to 40 years old) and a category of occupation. The written vignettes would be characteristic of one of the four MDT classifications identified by May and Rosedale (2012) as most prominent in patients with musculoskeletal knee pain: Derangement, Articular Dysfunction, Contractile Dysfunction and OTHER. Clinicians were asked to write the clinical vignettes on a blank McKenzie extremity assessment form (Appendix D). The blank McKenzie extremity assessment form used was revised from the standard form to exclude entry areas for patient names and other identifying information. To correspond with established prevalence, each clinician was asked to submit 6 clinical vignettes consisting of two Derangements, one Articular Dysfunction, one Contractile Dysfunction and one OTHER. A summary of these classifications is provided in Table 1.

Once received, all vignettes were reviewed by the author and a member of the advisory committee (SW & RR). The first reviewer (SW) is a MDT Credentialed physiotherapist and has 17 years of clinical practice working with patients with musculoskeletal knee pain. The second reviewer (RR) is a MDT Diploma physiotherapist with 23 years of clinical experience and is Senior Faculty of the McKenzie Institute. The review of the cases was done to ensure that each vignette was complete, possessed characteristics of one of the four MDT classifications requested and that a level of ambiguity existed that
would be present in the situation of a clinical patient presentation. Any discrepancies were identified, flagged and discussed with the subject who developed the vignette to ensure and verify accuracy of the case. In some situations, clinicians were unable to submit one or more of the number of vignettes matching the requested classification(s) because of the lack of a past patient assessment(s) that represented that classification(s). In those instances, clinicians chose another past patient assessment with one of the other requested classification(s) to submit to fulfill their quota of 6 clinical vignettes. In total, 60 vignettes were received: 24 Derangement, 8 Articular Dysfunction, 8 Contractile Dysfunction and 20 OTHER. After a review of all vignettes was completed, 53 cases were selected that were representative of the established prevalence for use in phase two of the study. Of the 53 vignettes, 22 were Derangement, 7 Articular Dysfunction, 7 Contractile Dysfunction and 17 OTHER. An example vignette for each category of classification can be found in Appendix E - H.

3.2.3 Phase 2

3.2.3.1 Participants

For phase two, six different raters were recruited based on previous willingness to participate in research from the publicly available list of MDT practitioners registered with McKenzie Institute International who practice in Canada or the United States. They were required to classify the 53 clinical vignettes generated in phase one of the study. The sample size of six raters was chosen to provide equal division of groups by level of MDT training and within group variability. To be included, the rater had to be a Credentialed or Diploma with the McKenzie Institute and have applied the MDT system to the extremities for more than three years. Raters were excluded if they participated in the creation of the clinical vignettes, did not wish to participate, were unable to understand written and spoken English, unable to provide informed consent or were unable to follow the instructions for rating the clinical vignettes. Correspondence was conducted and informed consent was obtained from each rater through electronic mail (Appendix I to J). In total, all six raters recruited consented to participation in this study.
3.2.3.2 Data Collection and Procedures

Demographic information like gender and age were collected along with other relevant characteristics like clinical practice setting, years of practice, length of time Credentialed/Diploma, discipline (eg. Physiotherapist versus Doctor of Chiropractic), proportion of extremity patients treated and proportion of knee patients treated with MDT. Data collection forms can be found in Appendix J. For each vignette, the raters were instructed to review the vignette and based on the history and clinical presentation, assign the vignette a classification of Derangement, Articular Dysfunction, Contractile Dysfunction or OTHER. Each vignette was randomly assigned a number from 1 to 53 to facilitate tracking of responses and data collection. All raters were blinded to the provisional MDT classification originally assigned to the vignette by its creator in phase one.

3.3 Analysis

Descriptive statistics for the demographic and clinical information for the raters were determined. Inter-rater reliability, our primary objective, was determined through the calculation of Fleiss kappa statistic along with 95% confidence interval (CI) and standard error (SE) across all six raters for all categories (Fleiss, 1971; Fleiss, Nee & Landis, 1979). Data were analyzed for Fleiss kappa using a program written in Matlab version 7.14 (Cardillo, 2007). Kappa values were interpreted using definitions outlined by Landis and Koch (1977): 0.01 to 0.20 slight agreement, 0.21 to 0.40 fair agreement, 0.41 to 0.60 moderate agreement, 0.61 to 0.80 substantial agreement and 0.81 to 1.00 almost perfect agreement.

It has been suggested that although overall kappa for three or more raters may lead to a better representation of reliability, overall kappa may mask extreme cases of agreement or disagreement for paired raters (O’Leary et al., 2014). A solution is to report both overall and paired kappa data to provide the most informative summary. Paired comparisons of the agreement of vignette classification among the six raters were thus analyzed and reported as percentage agreement and kappa statistic with standard error. Also, frequency distribution of the category of classification was analyzed for each
individual rater and reported as a whole number and percentage of total number of vignettes. Additionally, individual raters and their agreement with the vignette provisional classification assigned in phase one were analyzed and reported as percentage agreement and kappa statistic with standard error. Raters were grouped based on their level of education.

To examine if the level of education influenced the reliability, differences in Fleiss kappa values between Credentialed and Diploma therapists were compared. A bootstrap method with a 1000 samples was utilized and Fleiss kappa coefficients were calculated separately for the Credential and Diploma raters for each of these samples (McKenzie et al., 1996). The differences between the Fleiss kappa coefficients were determined. The mean of these differences was determined along with the 95% confidence interval represented by the 25 and 975 values. If the 95% confidence interval included zero, then no significant difference existed between the Credential and Diploma raters.

### 3.4 Results

The six raters recruited to rate the clinical vignettes were all physiotherapists and comprised of three Credentialed and three Diploma therapists. Four raters practiced fee for service and two worked in multiple settings. Four raters were male and two were female. Demographic information obtained from each of the raters is displayed in Table 3.

The overall kappa value amongst the six raters demonstrated substantial agreement with kappa=0.72 (SE=0.02) with a 95% CI of 0.71 to 0.73. The highest level of reliability was for the Derangement category with kappa=0.83; the lowest level of reliability was for the OTHER category with kappa=0.64. Articular and Contractile Dysfunction had a kappa of 0.67 and 0.69, respectively. There was 100% agreement in classification among all six raters in 31 of the 53 (58.5%) clinical vignettes.
### Table 3 Demographic information of participating phase 2 raters (n=6)

<table>
<thead>
<tr>
<th>Variables</th>
<th>Mean (SD)</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, years</td>
<td>51 (13.4)</td>
<td>35 to 67</td>
</tr>
<tr>
<td>Years of Practice</td>
<td>25 (13.7)</td>
<td>10 to 44</td>
</tr>
<tr>
<td>Proportion of Extremity Patients Treated in Practice, percentage</td>
<td>37 (16.0)</td>
<td>20 to 65%</td>
</tr>
<tr>
<td>Proportion of Knee Patients of Peripheral Joints in Practice, percentage</td>
<td>79 (37.2)</td>
<td>5 to 100%</td>
</tr>
</tbody>
</table>

SD - standard deviation
The frequency distribution of the category of classification by individual raters is displayed in Table 4. Derangement syndrome was the mostly commonly assigned classification to the vignettes across all raters, ranging from 20 (38%) – 26 (49%) of the total number (53) of vignettes reviewed.

Paired comparison of agreement in vignette classifications across the six raters are displayed in Table 5. The top right half shows percentage agreement and the bottom left half shows kappa scores (standard error) for all possible pairings of raters. The highest percentage of agreement (92%) and kappa (0.89) were between rater 1 and 5. The lowest percentage of agreement (72%) and kappa (0.58) were between rater 2 and 6. Reliability between raters showed moderate to substantial agreement.

Individual rater responses were compared to the provisional classifications of the vignettes and reliability calculated with results displayed in Table 6. Raters are grouped by level of MDT training. The highest percentage of agreement with the provisional classification was 91% for rater 5 and the lowest agreement was 81% for rater 6. Kappa scores ranged from 0.73 for rater 6 and 0.86 for rater 5. All raters showed substantial reliability (kappa ≥ 0.73) with the provisional classification.

Reliability by MDT training across raters for Credentialed and Diploma therapists are shown in Tables 7. The mean difference value between kappa values for Credentialed and Diploma therapists was -0.03 (95% CI -0.15 to 0.11). Since the confidence interval includes 0, there is no significant difference between rater groups based on level of education.
Table 4 Frequency distribution of category of classification by individual rater

<table>
<thead>
<tr>
<th>Rater</th>
<th>Classification n (%)</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Derangement</td>
<td>Articular Dysfunction</td>
</tr>
<tr>
<td>1</td>
<td>25 (47%)</td>
<td>6 (11%)</td>
</tr>
<tr>
<td>2</td>
<td>22 (41%)</td>
<td>6 (11%)</td>
</tr>
<tr>
<td>3</td>
<td>24 (45%)</td>
<td>7 (13%)</td>
</tr>
<tr>
<td>4</td>
<td>20 (38%)</td>
<td>11 (21%)</td>
</tr>
<tr>
<td>5</td>
<td>25 (47%)</td>
<td>7 (13%)</td>
</tr>
<tr>
<td>6</td>
<td>26 (49%)</td>
<td>5 (9%)</td>
</tr>
</tbody>
</table>

n - number of vignettes

Table 5 Percentage agreement and kappa (standard error) for paired comparisons among the six raters

<table>
<thead>
<tr>
<th>Percentage Agreement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rater</td>
</tr>
<tr>
<td>1</td>
</tr>
<tr>
<td>2</td>
</tr>
<tr>
<td>3</td>
</tr>
<tr>
<td>4</td>
</tr>
<tr>
<td>5</td>
</tr>
<tr>
<td>6</td>
</tr>
</tbody>
</table>

Kappa (Standard Error)
Table 6 Percentage agreement and kappa (standard error) of individual raters versus the provisional classification grouped by MDT education

<table>
<thead>
<tr>
<th>Rater</th>
<th>Statistic</th>
<th>Diploma</th>
<th>Diplomas</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>Provisional Classification</td>
<td>% Agreement</td>
<td>88%</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Kappa (SE)</td>
<td>0.83</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>(0.09)</td>
</tr>
</tbody>
</table>

SE – standard error

Table 7 Reliability by MDT education across raters

<table>
<thead>
<tr>
<th>MDT Education</th>
<th>Kappa (SE)</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Credentialed (n=3)</td>
<td>0.71 (0.048)</td>
<td>0.61 to 0.80</td>
</tr>
<tr>
<td>Diploma (n=3)</td>
<td>0.74 (0.051)</td>
<td>0.64 to 0.84</td>
</tr>
</tbody>
</table>

SE – standard error; CI – confidence interval
Chapter 4

4 Discussion

This chapter reviews the key findings of the research study and discusses the implications of these results for clinicians. Limitations of the study and recommendations for future research are also outlined.

4.1 Overview

The primary objective of this thesis was to determine the inter-rater reliability of the McKenzie System of MDT when trained therapists classify musculoskeletal knee pain using patient based clinical vignettes. The lack of research on the clinical utility of the MDT system when it is applied in the extremities, and specifically the knee, was the motivation for this thesis project.

4.2 Key Findings of the Thesis Project

The primary findings of this study suggest that the inter-rater reliability of Credentialed and Diploma clinicians within the MDT Institute demonstrate “substantial agreement” when using the MDT system to classify patients presenting with musculoskeletal knee pain (kappa=0.72). There was no statistically significant difference between Credentialed or Diploma raters (CI -0.15 to 0.11). Thus, it appears that clinicians with specific MDT training can use the MDT system to assess and classify patients with knee pain using clinical vignettes.

The results of this study are consistent with others evaluating the use of MDT in the extremity (Heidar Abady et al., 2014; Kelly, May, & Ross, 2008; May & Ross, 2009) and spine (Clare et al. 2005; Kilpikoski et al. 2002; Razmjou et al., 2000). The reliability of paired raters for the current study was kappa=0.58 to 0.89. Similarly, Razmajou et al. (2000) found overall reliability of kappa=0.70 when investigated the inter-rater reliability of two MDT trained examiners, one Diploma and one Credentialed, assessing real patients presenting with mechanical low back pain. Likewise, Kilpikoski et al. (2002) found an overall reliability between two MDT trained Diploma raters performing
independent, consecutive assessments of 39 patients with low back pain to be moderate (kappa=0.6). Using 14 raters, 7 Credentialed and 7 Diploma, Clare et al. (2005) reported the overall reliability for the classification of patients with lumbar and cervical pain was almost perfect (kappa=0.84) among paired raters. Hence, the MDT assessment appears to be a reliable method of assessment for patients presenting with musculoskeletal spinal or knee pain.

In the current study, individual raters demonstrated substantial reliability (kappa=0.73 to 0.86) while percentage of agreement ranged from 81 to 91% when rater classification was compared to the vignette provisional classification. Methodologically similar to the current study, Heidar Abady et al. (2014) used 54 clinical vignettes of patients presenting with musculoskeletal shoulder disorders to evaluate the inter-reliability of the MDT assessment by six Diploma clinicians and reported reliability of kappa=0.89 (0.77 to 0.96) and 95% overall agreement across raters against the provisional classification. Thus, MDT seems to have similar reliability when classifying clinical vignettes of patients presenting with musculoskeletal knee and shoulder pain.

The highest level of reliability was for the Derangement category with kappa=0.83 while the level of reliability for the three remaining categories varied with kappa=0.62 to 0.69. The difference between these levels of reliability may be explained to some degree by the general presentation of each category. The Derangement syndrome, by definition, is readily identifiable by a lasting reduction or elimination of patients’ symptoms through repeated movement in a particular direction (McKenzie and May, 2003). The relative lower level of agreement of the remaining categories may be partially attributable to the absence of a unique identifiable characteristic, such as directional preference that is present with Derangement syndrome. There may also be less familiarity with the extremity classifications and their criteria which have been more recently defined in the literature (May and Rosedale, 2012). This is particularly true for the OTHER category as multiple subgroups are included which makes determining a classification based exclusively on an initial assessment more challenging.
The relative reliability and similarity of kappa values between Credentialed (kappa=0.71) and Diploma (kappa=0.74) clinicians was anticipated. When grouped by education, Diploma raters compared to the provisional classification demonstrated reliability of kappa=0.78 to 0.83 and Credentialed raters reliability of kappa=0.73 to 0.86. Although Diploma holders of the MDT Institute undergo further education, each clinician in our study had a great deal of experience using the extremity assessment form and in treating patients with musculoskeletal pain in the extremities, and specifically, the knee. Thus, varying degrees of MDT competency did not appear to negatively impact the overall level of reliability of the system when evaluating musculoskeletal knee pain. However, we did not include raters without Credentialed or Diploma competence and as a result the effect of lower levels of MDT training was not evaluated.

4.3 Clinical Implications

It has been suggested that an assessment to simply identify structures as the cause of pain does not elicit enough information to understand the problem or to justify a course of management (Jones and Rivett, 2004). Indeed, specific features of the current model of examination for musculoskeletal knee pain, medical imaging and OSTs, have cast doubt on the clinical utility of this model. OSTs have demonstrated questionable diagnostic accuracy to discriminate the anatomical structures they are said to identify and medical imaging has brought to light the confounding prevalence of pathology and abnormalities in asymptomatic individuals. These findings would suggest a model less reliant on an anatomical diagnosis may be worth evaluating.

It has also been suggested that classification systems like MDT, may offer better clinical utility as the categories of classification are based on patient’s responses to repeated mechanical loading strategies rather than the presence of patho-anatomy (Lynch & May, 2013; May & Rosedale, 2012; Rosedale et al., 2014). An MDT assessment directs treatment with an appropriate loading strategy and in the presence of a directional preference, may determine who might and might not respond to treatment (Rosedale et al., 2014). An MDT assessment has shown to have good reliability in the spine and extremity, and while promising, more work needs to be done around the efficacy of MDT guided interventions.
There are two studies within the literature addressing inter-rater reliability of MDT in musculoskeletal pain in the lower extremity. (Kelly, May & Ross, 2008; May & Ross, 2009) To our knowledge, this is the first study to investigate the reliability of the MDT specifically for musculoskeletal knee pain alone. The results of this study on the knee reinforce the findings of previous reliability studies, indicating that the McKenzie System of MDT appears to be a reliable approach to assessing musculoskeletal pain in the knee.

Two previous studies have examined the use of an MDT guided intervention in the treatment of knee pain. Lynch & May (2013) documented the case study of directional preference of the knee using MDT. Although presenting with a positive McMurray’s test and pain with swimming, following prescribed exercise matching the directional preference, the patient reported 95% improvement in function and symptoms and a negative McMurray’s test. It was concluded by the authors that the use of McMurray’s test was not diagnostic and the result of the test appeared irrelevant and only clinical useful as a symptomatic baseline. More recently, Rosedale et al., (2014) published a randomized control trial using an MDT guided assessment to determine a directional preference for patients with end stage OA. Patients were classified as either Derangement or no Derangement. Patients who were matched with exercises consistent with a directional preference demonstrated significant decreases in pain and increases in self-report function scales after 2 weeks with large effect sizes (d=0.98 to 1.44). Although the effect sizes decreased at 3 months, they remained small to large (d=0.42 to 0.80) compared to the control group and patients without a directional preference. While the results from the study cannot be directly attributed to the MDT classification, it was concluded that the response to directional preference matched exercises should be explored further. With this in mind, it is reasonable to speculate that the method of MDT classification could facilitate the identification of who will or will not respond to treatment which in turn may enable the clinician to match the most appropriate treatment to various patient subgroups.

4.4 Limitations

Raters used in this study have achieved a high level of education and understanding of the McKenzie System of MDT and have significant experience applying the system to
musculoskeletal problems in the extremity. The background of the raters and subsequent findings will limit the generalizability of the results to those individuals with similar training and experience. As a result, generalizing the findings to practitioners without this level of training may not be appropriate. The vignettes with the provisional classification of OTHER were not further disseminated into subgroups for the raters to identify. To do so would have increased the number of categories of classification from 4 to 13, thus requiring an increase in the number of clinical vignettes for raters to review. As such, the reliability of raters classifying patients into those subgroups and the direction of subsequent treatment cannot be determined from this study. Additionally, the reviewers of each of the vignettes (SW, RR) were not blinded to the creator of the vignette or the provisional classification assigned to each vignette. This could result in the creation of a biased sample. Another potential limitation is the use of clinical vignettes as an alternative to real patients. It has been argued that vignettes cannot measure correspondence of hypothetical behavior and real world behavior (Evans et al., 2015) and may not capture the subtlety of a patients' presentation, oversimplify findings making a diagnosis easier and potentially inflating calculated agreement (Peabody et al., 2000; Werneke, Hart, Deutscher and Stratford, 2011). However, clinical vignettes offer the convenience of collecting information from multiple sources simultaneously while allowing for the flexibility of variable manipulation to ensure a heterogeneous sample.

4.5 Future Recommendations

This study found “substantial agreement” among Credentialed and Diploma holders in MDT. To generalize the use of the system to more users, future research should continue to investigate the reliability of MDT using raters with lower levels of training and experience.

Although the results of the study are encouraging, the results are limited to the reliability of raters classifying knee pain using clinical vignettes. To generalize the system further, future research should be conducted with real patients to demonstrate reliability in a real world, clinical setting.
Further to this, the efficacy of MDT guided treatment for patients presenting with musculoskeletal knee pain should be explored further. Long, Donelson & Fung (2004) found that exercises matching subjects’ directional preference in the lumbar spine significantly decreased pain and improved primary outcomes. The effect of directional preference matched exercises as indicated by the MDT classification needs to be evaluated and measured to substantiate use for clinical intervention in musculoskeletal knee pain.

4.6 Conclusion

In conclusion, this is the first study investigating the inter-rater reliability of the McKenzie System of MDT in the examination of musculoskeletal knee pain. The McKenzie System of MDT demonstrated substantial agreement, indicating acceptable inter-rater reliability for trained raters when using clinical vignettes to classify patients presenting with musculoskeletal knee pain. The results of this study offer preliminary support for the use of MDT in the assessment of musculoskeletal knee pain and support for future studies.


Appendices

Appendix A Ethics Approval from Western University Health Science Research Ethics Board

Western University Health Science Research Ethics Board
HSREB Delegated Initial Approval Notice

Principal Investigator: Dr. Trevor Birnbaum
Department & Institution: Health Science/Physical Therapy, Western University

HSREB File Number: 105498
Study Title: Intero-observer Reliability of the McKenzie System of Mechanical Diagnosis and Therapy in the Examination of the Knee

HSREB Initial Approval Date: August 26, 2014
HSREB Expiry Date: July 31, 2015

Documents Approved and/or Received for Information:

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<th>Document Name</th>
<th>Comments</th>
<th>Version Date</th>
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<td>2014.08.09</td>
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<td>Amendment</td>
<td>Amendment to Data Collection Form</td>
<td>2014.08.09</td>
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<tr>
<td>Amendment</td>
<td>Amendment to Lower Extremity Assessment Form</td>
<td>2014.08.09</td>
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<td>2014.08.09</td>
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<tr>
<td>Revised Letter of Information &amp; Consent Amendments to COH Phase 2</td>
<td>2014.08.09</td>
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The Western University Health Science Research Ethics Board (HSREB) has reviewed and approved the above named study, as of the HSREB Initial Approval Date noted above.

HSREB approval for this study remains valid until the HSREB Expiry Date noted above, conditional to timely submission and acceptance of HSREB Amendment/Delegation Review. If an Updated Approval Notice is required prior to the HSREB Expiry Date, the Principal Investigator is responsible for completing and submitting an HSREB Updated Approval Form in a timely fashion.

The Western University HSREB operates in compliance with the Tri-Council Policy Statement Ethical Conduct for Research Involving Human Subjects (TCP22), the International Conference on Harmonization of Technical Requirements for Registration of Pharmaceuticals for Human Use (ICH E6 R1), the Personal Health Information Protection Act (PHIPA, 2004), Part 4 of the Natural Health Product Regulations, Health Canada/Medical Device Regulations and Part C, Division 5, of the Food and Drug Regulations of Health Canada.

Members of the HSREB who are named as investigators in research studies do not participate in discussions related to, nor vote on such studies when they are presented to the HSREB.

The HSREB is registered with the U.S. Department of Health & Human Services under the IRB registration number IRB 00000946.
Appendix B Phase 1 Recruitment Email

Dear MDT Clinician,

You are invited to participate in a research study that is investigating the inter-examiner reliability of the McKenzie System of MDT when used by MDT trained clinicians in patients presenting with musculoskeletal knee pain. For this study, 10 clinicians are required to create 53 clinical vignettes for 6 expert MDT raters to review and classify.

We are asking you because you are Credentialed and/or have a Diploma standing within the McKenzie Institute and have been applying MDT to musculoskeletal problems in the extremity for greater than 3 years.

This study will be conducted by Trevor Birmingham, a Professor in the School of Physical Therapy at Western University in the School of Physical Therapy. Sean Willis, a Master of Science student in Health and Rehabilitation Sciences at Western University, Shawn Robbins, an Assistant Professor in the School of Physical and Occupational Therapy at McGill University and Richard Rosedale, an Instructor with the McKenzie Institute will also be participating in the study.

Attached is a Letter of Information for you to review and consider your participation. Also attached is a Consent Form for you to sign and submit should you wish to participate in this study.

Thank you for taking the time to consider participation in this study.

Sincerely,

Sean Willis
Appendix C Phase 1 Letter of Information and Consent Form

Letter of Information and Consent Form

Clinical Vignette Creation

Inter-observer Reliability of the McKenzie System of Mechanical Diagnosis and Therapy in the Examination of the Knee

Study Principal Investigator
Trevor Birmingham PhD, BSc(PT)
Western University, School of Physical Therapy

Study Co-Investigators
Sean Willis BSc(PT), MSc Student
Western University, Health and Rehabilitation Sciences

Shawn Robbins PhD, BSc(PT)
McGill University, School of Physical and Occupational Therapy

Study Advisor
Richard Rosedale PT, Dip MDT
Instructor, McKenzie Institute

1. Introduction

You are invited to participate in a research study that is investigating the inter-examiner reliability of the McKenzie System of Mechanical Diagnosis and Therapy (MDT) when used by MDT trained clinicians in patients presenting with musculoskeletal knee pain. For this study, 10 clinicians are required to create 53 clinical vignettes for 6 expert MDT raters to review and

Initials _______  LOI 08/08/2014
classify. The purpose of this letter is to provide you with the necessary background and
information to make an informed decision regarding your participation within this study.

2. Why Are We Asking You?

We are asking you because you are Credentialed and/or have a Diploma standing within the
McKenzie Institute and have been applying MDT to musculoskeletal problems in the extremity
for greater than 3 years.

We are giving this letter of information only to people who are registered on the publicly
available list of MDT clinicians practicing in Canada and the United States and who understand
both written and spoken English. If the above situation does not apply to you, we ask that you
do not volunteer to participate in our study.

3. Who is Conducting the Study?

This study will be conducted by Trevor Birmingham, a Professor in the School of Physical
Therapy at Western University in the School of Physical Therapy. Sean Willis, a Master of
Science student in Health and Rehabilitation Sciences at Western University, Shawn Robbins,
an Assistant Professor in the School of Physical and Occupational Therapy at McGill University
and Richard Rosedale, an Instructor with the McKenzie Institute will also be participating in the
study.
4. What will happen if you agree?

If you agree to participate, Sean Willis will contact you via email with a blank McKenzie Extremity Assessment form and instructions on generating clinical vignettes based on past patient assessment files. You will be asked to review your files and transfer the appropriate clinical information on the forms provided. This information should include a subjective history of the presenting musculoskeletal issue, signs and symptoms and relevant past medical history. The vignette should also include objective findings pertaining to baseline ROM, strength and neurological findings, functional baseline activities and responses to selected repeated movements. Each of the vignettes submitted are to be de-identified to only include age range (eg. 35-40 years old), gender and a category of occupation to respect the confidentiality of your patients. You are asked to provide clinical vignettes that are characteristic of the following four mechanical syndromes: (2) derangement, (2) other, (1) contractile dysfunction and (1) articular dysfunction.

It is expected that the review of your past patient assessments and creation of the vignettes will take some time. The anticipated time commitment for the creation of the clinical vignettes is 1 hour per vignette for a total of 6 hours. 3 months will be provided to complete this process.
5. What are the risks and benefits?

There are no direct benefits from participating in this study. However, for clinicians who treat people with musculoskeletal knee disorders, this would be the first study to address inter-examiner reliability of McKenzie system in assessing patients with knee pain. It would provide MDT practitioners with valuable information about reliability of this assessment method.

There are no risks to participating in this study.

6. What about confidentiality?

Your confidentiality will be protected. No information that discloses your identity will be released or be a part of any publication of the results of the study. Your first and last name as well as your email address will be collected for contact and communication purposes only. All records will be given a code ID to be used on all collected forms. All information collected will be kept in locked filing cabinets and shredded seven years after the completion of the study.

7. Alternatives to participation?

Participation in this study is voluntary. You may refuse to participate, refuse to answer any questions or withdraw from the study at any time with no effect on your future employment. Should you withdraw from the study, you may also withdraw any vignettes that you have submitted.
8. If you agree to participate in the study

Please sign the attached consent form and return to Sean Willis. You do not waive any of your legal rights by signing the consent form.

You may keep this letter of information. A copy of your signed consent form will be made for you. If you have any questions about this study, please contact Sean Willis.

If you have any questions about your rights as a research participant or the conduct of the study, you may contact The Office of Research Ethics at Western University.

Representatives of The University of Western Ontario Health Sciences Research Ethics Board may contact you or require access to your study-related records to monitor the conduct of the research.
Consent Form

Inter-observer Reliability of the McKenzie System of Mechanical Diagnosis and Therapy in the Examination of the Knee

I have read the accompanying letter of information and I agree to participate. Questions I had have been answered to my satisfaction.

<table>
<thead>
<tr>
<th>Date</th>
<th>Participant’s Name</th>
<th>Participant’s Signature</th>
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<th>Name of Person Obtaining Informed Consent</th>
<th>Signature of Person Obtaining Informed Consent</th>
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Initials _______  LOI 08/08/2014
Appendix D McKenzie Institute Lower Extremities Assessment Form

THE MCKENZIE INSTITUTE
LOWER EXTREMITIES ASSESSMENT

Vignette ID #: __________

Age Range (eg. 35-45 yrs.) : __________________
Gender : M F
Category of Occupation : __________________________

Work: Mechanical stresses
Leisure: Mechanical stresses
Functional disability from present episode

Functional disability score
VAS Score (0-10)

HISTORY

Present symptoms
Present since
Commenced as a result of
Symptoms at onset
Spinal history
Constant symptoms: Intramuscular symptoms:

Worse
bending sitting / rising / first few steps standing walking stairs squatting / kneeling
am / as the day progresses / pm when still / on the move Sleeping: prone / sup / side R / L
Other

Better
bending sitting standing walking stairs squatting / kneeling
am / as the day progresses / pm when still / on the move Sleeping: prone / sup / side R / L
Other

Continued use makes the pain: Better Worse No Effect Disturbed night Yes / No

Pain at rest Yes / No
Site: Back / Hip / Knee / Ankle / Foot

Other Questions: Swelling Clicking / Locking Giving Way / Falling

Previous episodes
Previous treatments
General health: Good / Fair / Poor

Medications: Nil / NSAIDS / Analg / Steroids / Anticoag / Other
Imaging: Yes / No

Recent or major surgery: Yes / No
Night pain: Yes / No

Accidents: Yes / No
Unexplained weight loss: Yes / No

Summary
Acute / Sub-acute / Chronic

Trauma / Insidious Onset

Sites for physical examination
Back / Hip / Knee / Ankle / Foot

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Version 04/08/2014
EXAMINATION

POSTURE
Sitting: Good / Fair / Poor
Correction of Posture: Better / Worse / No Effect / NA
Standing: Good / Fair / Poor
Other observations:

NEUROLOGICAL: NA / Motor / Sensory / Reflexes / Dural

BASELINES (pain or functional activity):

EXTREMITIES
Hip / Knee / Ankle / Foot

<table>
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<tr>
<th>MOVEMENT LOSS</th>
<th>Maj</th>
<th>Mod</th>
<th>Min</th>
<th>Nil</th>
<th>Pain</th>
<th>Maj</th>
<th>Mod</th>
<th>Min</th>
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</table>

Passive Movement (+/- over pressure) (note symptoms and range):
PDM ERP

Resisted Test Response (pain)

Other Tests

SPINE
Movement Loss
Effect of repeated movements
Effect of static positioning
Spine testing: Not relevant / Relevant / Secondary problem

Baseline Symptoms

<table>
<thead>
<tr>
<th>Repeated Tests</th>
<th>Symptom Response</th>
<th>Mechanical Response</th>
</tr>
</thead>
<tbody>
<tr>
<td>Active/Passive movement, resisted test, functional test</td>
<td>During = Produce, Abolish, Increase, Decrease, NE</td>
<td>After = Better, Worse, NB, NW, NE</td>
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<td>Effect = ↑ or ↓ ROM, strength or key functional test</td>
<td>No Effect</td>
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<tr>
<td>Effect of static positioning</td>
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PROVISIONAL CLASSIFICATION
Extremities Spine
Dysfunction - Articular Contractile
Displacement Postural
Other Uncertain

PRINCIPLE OF MANAGEMENT
Education
Exercise and Dosage
Treatment Goals

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Version 04/08/2014
Appendix E Derangement Vignette

THE McKENZIE INSTITUTE
LOWER EXTREMITIES ASSESSMENT

Vignette ID #: 2

Age Range (eg. 35-45 yrs.): 65-75
Gender: M F
Category of Occupation: Admin Assistant

Work: Mechanical stresses ___up and down, squat, kneeling___
Leisure: Mechanical stresses ____________________________
Functional disability from present episode: Kneeling
Functional disability score ____________________________
VAS Score (0-10) ____________________________

HISTORY

Present symptoms:
Present since: 3 Weeks Improving / Unchanging / Worsening
Commenced as a result of: Lots of hiking, walking and knee gave way Or No Apparent Reason
Symptoms at onset: Sharp knee pain Paraesthesia: Yes / No
Spinal history: Denies Cough / Sneeze +ve / -ve

Constant symptoms: Intermittent Symptoms: Knee pain
Worse: bending sitting / rising / first few steps standing walking stairs squatting / kneeling
am / as the day progresses / pm when still / on the move Sleeping: prone / sup / side R / L
Other: Rowing machine, treadmill fast
Better: bending sitting standing walking stairs squatting / kneeling
am / as the day progresses / pm when still / on the move Sleeping: prone / sup / side R / L
Other: Lying with leg straight

Continued use makes the pain: Better Worse No Effect Disturbed night Yes / No
Pain at rest: Yes / No Site: Back / Hip / Knee / Ankle / Foot
Other Questions: Swelling Clicking / Locking Giving Way / Falling

Previous episodes: A couple of years ago - resolved
Previous treatments: None
General health: Good / Fair / Poor
Medications: Nil / NSAIDS / Analg / Steroids / Anticoag / Other Ibuprofen
Imaging: Yes / No X-ray – OA changes
Recent or major surgery: Yes / No Night pain: Yes / No
Accidents: Yes / No Unexplained weight loss: Yes / No

Summary: Acute / Sub-acute / Chronic Trauma / Insidious Onset
Sites for physical examination: Back / Hip / Knee / Ankle / Foot Other: ______

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EXAMINATION

POSTURE
Sitting: Good / Fair / Poor
Standing: Good / Fair / Poor
Other observations:

NEUROLOGICAL:
NA / Motor / Sensory / Reflexes / Dural

BASELINES (pain or functional activity):

EXTREMITIES
Hip / Knee / Ankle / Foot

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Adduction / Inversion
Abduction / Eversion
Internal Rotation
External Rotation

Passive Movement (+/- over pressure) (note symptoms and range):
As active movements

Resisted Test Response (pain)
Not tested

Other Tests

SPINE
Movement Loss
Effect of repeated movements
Effect of static positioning
Spine testing: Not relevant / Relevant / Secondary problem

Baseline Symptoms

<table>
<thead>
<tr>
<th>Repeated Tests</th>
<th>Symptom Response</th>
<th>Mechanical Response</th>
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</thead>
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<td>After = Better, Worse, NB, NW, NE</td>
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<td>Rep Extension</td>
<td>Produce ERP</td>
<td>NW</td>
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<td>Rep OKC Extension</td>
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<td>NW</td>
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<td>Produce ERP, less with repetition</td>
<td>Increase flexion</td>
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<td>Produce ERP</td>
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Effect of static positioning

PROVISIONAL CLASSIFICATION
Extremities
Spine
Dysfunction – Articular
Dysfunction – Contractile
Derangement
Postural
Other
Uncertain

PRINCIPLE OF MANAGEMENT
Education
Exercise and Dosage
Treatment Goals

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Appendix F Articular Dysfunction Vignette

THE McKENZIE INSTITUTE
LOWER EXTREMITIES ASSESSMENT

Vignette ID #: 1

Age Range (eg. 35-45 yrs.): 45-55
Gender: M F
Category of Occupation: 

Work: Mechanical stresses Sitting
Leisure: Mechanical stresses 
Functional disability from present episode
Functional disability score
VAS Score (0-10)

Present symptoms Knee pain
Present since 18 months Improving / Unchanging / Worsening
Commenced as a result of Twisted knee Or No Apparent Reason
Symptoms at onset Knee pain Parasthesia: Yes / No
Spinal history Cough / Sneeze +ve / -ve
Constant symptoms: Intermittent Symptoms: Knee Pain
Worse bending sitting / rising / first few steps standing walking stairs squattting / kneeling
em / as the day progresses / pm when still / on the move Sleeping: prone / sup / side R / L
Other
Better bending sitting standing walking stairs squattting / kneeling
em / as the day progresses / pm when still / on the move Sleeping: prone / sup / side R / L
other
Continued use makes the pain: Better Worse No Effect Disturbed night Yes / No
Pain at rest Yes / No Site: Back / Hip / Knee / Ankle / Foot
Other Questions: Swelling Clicking / Locking Giving Way / Falling

Previous episodes 
Previous treatments 
General health: Good / Fair / Poor
Medications: Nil / NSAIDS / Analg / Steroids / Antiocoag / Other HTN meds
Imaging: Yes / No X-ray - DJD
Recent or major surgery: Yes / No Meniscectomy 16 months ago Night pain: Yes / No
Accidents: Yes / No Unexplained weight loss: Yes / No
Summary Acute / Sub-acute / Chronic Trauma / Insidious Onset
Sites for physical examination Back / Hip / Knee / Ankle / Foot Other: 

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**EXAMINATION**

**POSTURE**
- Sitting: Good / Fair / Poor
- Correction of Posture: Better / Worse / No Effect / NA
- Standing: Good / Fair / Poor
- Other observations: 

**NEUROLOGICAL:**  NA / Motor / Sensory / Reflexes / Dural

**BASELINES (pain or functional activity):** Squat, step down

**EXTREMITIES**  Hip / Knee / Ankle / Foot

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<th>Mod</th>
<th>Min</th>
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**Passive Movement (+/- over pressure) (note symptoms and range):**
- Extension – full with over pressure
- Flexion – moderate loss

**Resisted Test Response (pain):** No pain with resisted flexion or extension

**Other Tests:** 

**SPINE**

- Movement Loss
- Effect of repeated movements
- Effect of static positioning
- Spine testing: Not relevant / Relevant / Secondary problem

**Baseline Symptoms**

<table>
<thead>
<tr>
<th>Repeated Tests</th>
<th>Symptom Response</th>
<th>Mechanical Response</th>
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</thead>
<tbody>
<tr>
<td>Active/Passive movement, resisted test, functional test</td>
<td>During – Produce, Abolish, Increase, Decrease, NE</td>
<td>After – Better, Worse, NB, NW, NE</td>
</tr>
<tr>
<td>Rep passive flexion</td>
<td>Produce</td>
<td>NW</td>
</tr>
<tr>
<td>Rep loaded flexion</td>
<td>Produce</td>
<td>NW</td>
</tr>
<tr>
<td>Rep active extension</td>
<td>NE</td>
<td>NE</td>
</tr>
<tr>
<td>Effect of static positioning</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**PROVISIONAL CLASSIFICATION**

- Extremities: Contractile
- Spine: Postural

**PRINCIPLE OF MANAGEMENT**

- Education
- Exercise and Dosage
- Treatment Goals

McKenzie Institute International 2014
Version 04/08/2014
Appendix G Contractile Dysfunction Vignette

THE MCKENZIE INSTITUTE
LOWER EXTREMITIES ASSESSMENT

Vignette ID #: ______

Age Range (eg. 35-45 yrs.): ______ 10-20 ______

Gender: M F

Category of Occupation: ______ Student ______

Work: Mechanical stresses ______ School ______

Leisure: Mechanical stresses ______ Football - Punter ______

Functional disability from present episode: ______ Pain with kicking ______

Functional disability score

VAS Score (0-10)

HISTORY

Present symptoms: ______ Left anterior knee pain ______

Present since: 3 weeks

Commenced as a result of: ______ With kicking ______

Symptoms at onset: ______ Anterior knee pain ______

Spinal history: ______ Denies ______

Constant symptoms: ______ Intermitent symptoms: ______ Anterior knee pain ______

Worse

bending sitting / rising / first few steps standing walking stairs kneeling

am / as the day progresses / pm when still / on the move Sleeping: prone / sup / side R/L

Better

bending sitting standing walking stairs squatting / kneeling

am / as the day progresses / pm when still / on the move Sleeping: prone / sup / side R/L

other ______

Continued use makes the pain: ______ Better ______ Worse ______ No Effect ______

Disturbed night: Yes / No

Pain at rest: Yes – after kicking / No ______

Site: Back / Hip / Knee / Ankle / Foot ______

Other Questions: ______ Swelling ______ Clicking / Locking ______

Giving Way / Failing ______

Previous episodes: ______ Last football season ______

Previous treatments: ______ None ______

General health: ______ Good / Fair / Poor ______

Medications: ______ Nil / NSAIDS / Analg / Steroids / Anticoag / Other ______

Imaging: ______ Yes / No ______

Recent or major surgery: ______ Yes / No ______

Night pain: Yes / No ______

Accidents: ______ Yes / No ______

Unexplained weight loss: ______ Yes / No ______

Summary: ______ Acute / Sub-acute / Chronic ______

Trauma / Insidious Onset ______

Sites for physical examination: ______ Back / Hip / Knee / Ankle / Foot ______

Other: ______

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EXAMINATION

POSTURE
Sitting: Good / Fair / Poor
Correction of Posture: Better / Worse / No Effect / NA
Standing: Good / Fair / Poor
Other observations:

NEUROLOGICAL:
NA / Motor / Sensory / Reflexes / Dural

BASELINES (pain or functional activity):
Squatting

EXTREMITIES
Hip / Knee / Ankle / Foot

<table>
<thead>
<tr>
<th>MOVEMENT LOSS</th>
<th>Meq</th>
<th>Mod</th>
<th>Min</th>
<th>Nil</th>
<th>Pain</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flexion</td>
<td></td>
<td></td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Extension</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dorsal Flexion</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Plantar Flexion</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Adduction / Inversion
Abduction / Eversion
Internal Rotation
External Rotation

Passive Movement (+/- over pressure) (note symptoms and range):
PDM ERP
No loss of extension with over pressure
No loss of flexion with over pressure

Resisted Test Response (pain):
Pain with resisted extension
Pain free resisted flexion

Other Tests

SPINE
Movement Loss
Effect of repeated movements
Effect of static positioning
Spine testing: Not relevant / Relevant / Secondary problem

Baseline Symptoms:
None

<table>
<thead>
<tr>
<th>Repeated Tests</th>
<th>Symptom Response</th>
<th>Mechanical Response</th>
</tr>
</thead>
<tbody>
<tr>
<td>Active/Passive movement, resisted test, functional test</td>
<td>During — Produce, Abolish, Increase, Decrease, NE</td>
<td>After — Better, Worse, NB, NW, NE</td>
</tr>
<tr>
<td>Rep. Passive Extension</td>
<td>NE</td>
<td>NE</td>
</tr>
<tr>
<td>Rep. Active Extension</td>
<td>Produce</td>
<td>NW</td>
</tr>
<tr>
<td>Rep. Flexion with OP</td>
<td>Produce</td>
<td>NW</td>
</tr>
</tbody>
</table>

Effect of static positioning

PROVISIONAL CLASSIFICATION
Extremities
Spine
Dysfunction — Articular
Derangement
Other
Contractile
Postural
Uncertain

PRINCIPLE OF MANAGEMENT
Education
Exercise and Dosage
Treatment Goals

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Version 04/08/2014
Appendix H OTHER Vignette

**THE McKENZIE INSTITUTE**  
**LOWER EXTREMITIES ASSESSMENT**

| Vignette ID #: 10 |

---

**Age Range (eg. 35-45 yrs.):** 15-25

**Gender:** M F

**Category of Occupation:** Student

**Work:** Mechanical stresses

**Leisure:** Mechanical stresses Soccer

Functional disability from present episode

Functional disability score

VAS Score (0-10)

<table>
<thead>
<tr>
<th>Present symptoms</th>
<th>Knee pain</th>
</tr>
</thead>
<tbody>
<tr>
<td>Present since</td>
<td>2 weeks</td>
</tr>
<tr>
<td>Commenced as a result of</td>
<td>Twist playing soccer</td>
</tr>
<tr>
<td>Symptoms at onset</td>
<td>Cough / Sneeze +ve / -ve</td>
</tr>
<tr>
<td>Spinal history</td>
<td>Paraesthesia: Yes / No</td>
</tr>
<tr>
<td>Constant symptoms:</td>
<td>Intermittent Symptoms: Knee Pain</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Worse</th>
<th>bending sitting / rising / first few steps standing walking stairs squatting / kneeling am / as the day progresses / pm when still / on the move</th>
</tr>
</thead>
<tbody>
<tr>
<td>Other</td>
<td>Sleeping: prone / sup / side R / L</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Better</th>
<th>bending sitting standing walking stairs squatting / kneeling am / as the day progresses / pm when still / on the move</th>
</tr>
</thead>
<tbody>
<tr>
<td>Other</td>
<td>Sleeping: prone / sup / side R / L</td>
</tr>
</tbody>
</table>

**Continued use makes the pain:** Better Worse No Effect

**Pain at rest:** Yes / No

**Site:** Back / Hip / Knee / Ankle / Foot

**Other Questions:** Swelling Clicking / Locking Giving Way / Failing

**Previous episodes**

**Previous treatments** Ice

**General health:** Good / Fair / Poor

**Medications:** Nil / NSAIDS / Analg / Steroids / Anticoag / Other

**Imaging:** Yes / No

**Recent or major surgery:** Yes / No

**Night pain:** Yes / No

**Night pain:** Yes / No

**Accidents:** Yes / No

**Unexplained weight loss:** Yes / No

**Summary**

<table>
<thead>
<tr>
<th>Acute / Sub-acute / Chronic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trauma / Insidious Onset</td>
</tr>
</tbody>
</table>

**Sites for physical examination**

<table>
<thead>
<tr>
<th>Back / Hip / Knee / Ankle / Foot</th>
</tr>
</thead>
</table>

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EXAMINATION

POSTURE
Sitting: Good / Fair / Poor
Standing: Good / Fair / Poor
Other observations:

NEUROLOGICAL: NA / Motor / Sensory / Reflexes / Dural

BASELINES (pain or functional activity): Squat, steps

EXTREMITIES
Hip / Knee / Ankle / Foot

<table>
<thead>
<tr>
<th>MOVEMENT LOSS</th>
<th>Maj</th>
<th>Mod</th>
<th>Min</th>
<th>Nil</th>
<th>Pain</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flexion</td>
<td></td>
<td></td>
<td>X</td>
<td>ERP</td>
<td>Adduction / Inversion</td>
</tr>
<tr>
<td>Extension</td>
<td></td>
<td>X</td>
<td>ERP</td>
<td></td>
<td>Abduction / Eversion</td>
</tr>
<tr>
<td>Dorsi Flexion</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Internal Rotation</td>
</tr>
<tr>
<td>Plantar Flexion</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>External Rotation</td>
</tr>
</tbody>
</table>

Passive Movement (+/- over pressure) (note symptoms and range):
Flexion and extension WNL

Resisted Test Response (pain) Quads and hams 4+/5

Other Tests

SPINE
Movement Loss
Effect of repeated movements
Effect of static positioning
Spine testing: Not relevant / Relevant / Secondary problem

Baseline Symptoms

<table>
<thead>
<tr>
<th>Repeated Tests</th>
<th>Symptom Response</th>
<th>Mechanical Response</th>
</tr>
</thead>
<tbody>
<tr>
<td>Active/Passive movement, resisted test, functional test</td>
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</tr>
<tr>
<td>Rep flexion</td>
<td>Produce</td>
<td>NW</td>
</tr>
<tr>
<td>Rep extension</td>
<td>Produce</td>
<td>NW</td>
</tr>
<tr>
<td>Effect of static positioning</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

PROVISIONAL CLASSIFICATION
Extremities: Spine
Dysfunction - Articular Contractile
Displacement Postural
Other Uncertain

PRINCIPLE OF MANAGEMENT
Education
Exercise and Dosage
Treatment Goals

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Appendix I Phase 2 Recruitment Email

Dear MDT Clinician,

You are invited to participate in a research study that is investigating the inter-examiner reliability of the McKenzie System of MDT when used by MDT trained clinicians in patients presenting with musculoskeletal knee pain. For this study, 6 expert MDT raters are required to review and classify 53 clinical vignettes.

We are asking you because you are Credentialed and/or have a Diploma standing within the McKenzie Institute and have been applying MDT to musculoskeletal problems in the extremity for greater than 3 years.

This study will be conducted by Trevor Birmingham, a Professor in the School of Physical Therapy at Western University in the School of Physical Therapy. Sean Willis, a Master of Science student in Health and Rehabilitation Sciences at Western University, Shawn Robbins, an Assistant Professor in the School of Physical and Occupational Therapy at McGill University and Richard Rosedale, an Instructor with the McKenzie Institute will also be participating in the study.

Attached is a Letter of Information for you to review and consider your participation. Also attached is a Consent Form for you to sign and submit should you wish to participate in this study.

Thank you for taking the time to consider participation in this study.

Sincerely,

Sean Willis
Appendix J Phase 2 Letter of Information and Consent Form

Western HealthSciences

Letter of Information and Consent Form

Classification of Clinical Vignettes

Inter-observer Reliability of the McKenzie System of Mechanical Diagnosis and Therapy in the Examination of the Knee

Study Principal Investigator
Trevor Birmingham PhD, BSc(PT)
Western University, School of Physical Therapy

Study Co-Investigators
Sean Willis BSc(PT), MSc Student
Western University, Health and Rehabilitation Sciences
Shawn Robbins PhD, BSc(PT)
McGill University, School of Physical and Occupational Therapy

Study Advisor
Richard Rosedale PT, Dip MDT
Instructor, McKenzie Institute

1. Introduction

You are invited to participate in a research study that is investigating the inter-examiner reliability of the McKenzie System of MDT when used by MDT trained clinicians in patients presenting with musculoskeletal knee pain. For this study, 6 expert MDT raters are required to

Initials________  LOI 08/08/2014
review and classify 53 clinical vignettes. The purpose of this letter is to provide you with the necessary background and information to make an informed decision regarding your participation within this study.

2. Why Are We Asking You?

We are asking you because you are Credentialed and/or have a Diploma standing within the McKenzie Institute and have been applying MDT to musculoskeletal problems in the extremity for greater than 3 years.

We are giving this letter of information only to people who are registered on the publically available list of MDT clinicians practicing in Canada and the United States, who understand both written and spoken English and who did not participate in generating the 53 clinical vignettes in the study. If the above situation does not apply to you, we ask that you do not volunteer to participate in our study.

3. Who is Conducting the Study?

This study will be conducted by Trevor Birmingham, a Professor in the School of Physical Therapy at Western University in the School of Physical Therapy. Sean Willis, a Master of Science student in Health and Rehabilitation Sciences at Western University, Shawn Robbins, an Assistant Professor in the School of Physical and Occupational Therapy at McGill University and

Initials_______

LOI 08/28/2014

2
Richard Rosedale, an Instructor with the McKenzie Institute will also be participating in the study.

4. What will happen if you agree?

If you agree to participate, Sean Willis will contact you via email with 53 clinical vignettes and instructions regarding classifying each of the vignettes into one of four mechanical syndromes: derangement, articular dysfunction, contractile dysfunction and other.

In addition to completing the classification of the clinical vignettes, you will also be asked to complete and submit a data collection form to Sean Willis that will record your: gender, age, country of practice, clinical practice setting, years of practice, length of time credentialed/degree, discipline (e.g. PT versus DC), proportion of extremity patients treated and proportion of knee patients treated with MDT. This information will be used within the study to describe the clinical background and characteristics of the group of expert raters as well as analyze the effect those characteristics may have on the level of agreement between raters.

It is expected that the review and subsequent rating of the 53 clinical vignettes may take some time. The anticipated time commitment to review and rate each clinical vignette is 30 minutes for a total of 27 hours to rate all vignettes. A total of 2 months will be provided for this process.

Initials _______  LOI 08/08/2014
5. What are the risks and benefits?

There are no direct benefits from participating in this study. However, for clinicians who treat people with musculoskeletal knee disorders, this would be the first study to address inter-examiner reliability of Mckenzie system in assessing patients with knee pain. It would provide MDT practitioners with valuable information about reliability of this assessment method.

There are no risks to participating in this study.

6. What about confidentiality?

Your confidentiality will be protected. No information that discloses your identity will be released or be a part of any publication of the results of the study. Your first and last name as well as your address, email address and phone number will be collected for contact and communication purposes only. All records will be given a code ID to be used on all collected forms. All information collected will be kept in locked filing cabinets and shredded seven years after the completion of the study.

7. Alternatives to participation?

Participation in this study is voluntary. You may refuse to participate, refuse to answer any questions or withdraw from the study at any time with no effect on your future employment.

Initials ____________________________  LOI 08/08/2014
Should you withdraw from the study, you may also withdraw any information or data that you have submitted.

8. If you agree to participate in the study

Please sign the attached consent form and return to Sean Willis by email or fax. You do not waive any of your legal rights by signing the consent form.

You may keep this letter of information. A copy of your signed consent form will be made for you. If you have any questions about this study, please contact Sean Willis.

If you have any questions about your rights as a research participant or the conduct of the study, you may contact The Office of Research Ethics at Western University.

Representatives of The University of Western Ontario Health Sciences Research Ethics Board may contact you or require access to your study-related records to monitor the conduct of the research.
Consent Form

Inter-observer Reliability of the McKenzie System of Mechanical Diagnosis and Therapy in the Examination of the Knee

I have read the accompanying letter of information and I agree to participate. Questions I had have been answered to my satisfaction.

<table>
<thead>
<tr>
<th>Date</th>
<th>Participant's Name</th>
<th>Participant's Signature</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Date</th>
<th>Name of Person Obtaining Informed Consent</th>
<th>Signature of Person Obtaining Informed Consent</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Initials

LOI 08/08/2014
Appendix K Phase 2 Data Collection Form

Subject Characteristics

Age: ___________ Years of Practice: ________________ Diploma/Credentialed

Clinical Practice Setting: Private Practice
Insurance/Workman’s Compensation
Hospital Outpatient Setting
Physician Referral

Discipline: Physical Therapist

Chiropractor

Proportion of Caseload that are Extremity Patients: ____________________________

Proportion of Knee patients treated with MDT: ________________________________

Contact Information

Name (Please Print): ____________________________

Address: ____________________________

City: ____________________________

Email: ____________________________

Phone Number (H): ____________________________ (W): ____________________________
Curriculum Vitae

Name: Sean Willis

**Post-secondary Education and Degrees:**
- The University of Western Ontario, London, Ontario, Canada
- 1994 - 1998 BSc (PT)

**Post-graduate Qualifications:**
- MDT Credentialing Exam
- April 2008

**Related Work Experience:**
- Physiotherapist
  - London Health Sciences Centre
  - 2000 - Present

  - Clinical Associate
    - Western University, School of Physical Therapy
    - 2005 - Present

  - Teaching Assistant
    - Western University
    - 2015