COMBINING ORTHOPEDIC SPECIAL TESTS TO IMPROVE DIAGNOSIS OF SHOULDER PATHOLOGY

Authors:
Corresponding Author
Eric J, Hegedus PT, DPT, MHSc
Professor and Chair, High Point University, Department of Physical Therapy, High Point NC 27262
ehegedus@highpoint.edu
336-906-2133

Chad Cook PT, PhD, MBA, FAAOMPT
Professor, Duke University, Doctor of Physical Therapy Program, Duke University, Durham, NC

Dr Jeremy Lewis FCSP MMACP, Reader in Physiotherapy, University of Hertfordshire Department of Allied Health Professions and Midwifery, School of Health and Social Work

Dr Alexis Wright PT, PhD, FAAOMPT
Assistant Professor, High Point University, Department of Physical Therapy, High Point, NC

Jin-Young Park, MD
Professor, Shoulder, Elbow & Sports Center, Konkuk University, Seoul, South Korea
ABSTRACT

The use of orthopedic special tests (OSTs) to diagnose shoulder pathology via the clinical examination is standard in clinical practice. There is a great deal of research on special tests but much of the research is of a lower quality implying that the metrics from that research, sensitivity, specificity, and likelihood ratios, is likely to vary greatly in the hands of different clinicians and in varying practice environments. A way to improve the clinical diagnostic process is to cluster OSTs and to use these clusters to either rule in or out different pathologies. The aim of the article is to review the best OST clusters, examine the methodology by which they were derived, and illustrate, with a case study, the use of these OST clusters to arrive at a pathology-based diagnosis.

Key Words: Likelihood ratios, Shoulder, Diagnosis
INTRODUCTION

Physical examination of the shoulder involves a series of steps typically beginning with history, progressing with motion and muscle testing, and culminating in the use of orthopedic special tests (OSTs) with the aim of diagnosing shoulder pathology. While the process itself is systematic and straightforward, for evidence-based practitioners, there are numerous problems encountered when trying to arrive at a diagnosis. First, there is little evidence reporting the diagnostic accuracy of critical pieces of the clinical examination such as history, motion testing, and muscle testing causing a greater reliance on OSTs. Second, although there is a great deal of research on OSTs of the shoulder, much of that research is of moderate to low quality (Hegedus, et al., 2008; Hegedus, et al., 2012). Third, even in those OSTs that come from high quality literature, there are very few that display solid diagnostic metrics, high sensitivity and specificity (Hegedus, et al., 2008; Hegedus, et al., 2012). Fourth, although sensitivity and specificity are helpful internal test metrics, there are issues in the application of these metrics to clinical practice. Finally, clinicians and researchers improve diagnostic accuracy by clustering OSTs together; however, in some cases, the clusters are used incorrectly or provide metrics that lead to post-test probabilities that are no different than use of a single stand alone test.

Our aims in this paper are to discuss the importance of likelihood ratios and modified probability in the diagnostic process, to explain multivariate modeling and outline the most effective methods to combine tests for either screening or confirmation of diagnosis. For context, we'll briefly review the best test clusters that
have been published, and finally, we'll use a case study to illustrate how the best available test clusters should be used to aid in diagnosis.

**LIKELIHOOD RATIOS AND MODIFIED PROBABILITY**

Diagnostic accuracy studies have design consistencies, standardized metrics, and assumptions. First and foremost, all diagnostic accuracy studies enroll populations of individuals with and without the condition of interest; the condition of interest being the diagnosis studied. Those without the condition of interest should be individuals with some other competing health malady that would normally be distinguished in a traditional clinical environment.

The simplest measures of diagnostic accuracy are sensitivity and specificity. Sensitivity is the proportion of people *with the condition of interest* who will have a *positive* result, whereas specificity is the proportion of the patients *do not have the condition of interest* who have a *negative* result. Mathematically, sensitivity values are calculated only from those with the condition of interest, whereas specificity values are calculated from those without the condition of interest. This is one reason that the use of these internal metrics is limited. For example, sensitivity fails to recognize any of the examination findings that are reflective of those who did not have the condition of interest.

Both sensitivity and specificity are reported in percentages, from 0 to 100. A 100% sensitivity or specificity suggests that the test will be positive 100% of the time (if truly sensitivity) in patients with the condition of interest and will be appropriately negative in 100% of cases when the patient does not have the condition of interest. In order to emphasize the context associated with these measures, a sensitivity of 20%,
which is a finding associated with most reflex testing suggests that the test will be positive in only 20% of cases in which the patient actually has the condition of interest.

Likelihood ratios and probability metrics are calculated from the sensitivity and specificity values. Positive and negative predictive values reflect proportions of positive and negative results when the clinical tests are employed. Positive predictive value (PPV), which is sometimes referred to as precision, is the number of true positives divided by the number of true positive plus the number of false positives. The calculation is derived from positive finding from those with and without the condition of interest. A negative predictive value (NPV) is the number of true negative divided by the number of true negatives plus the number of false negatives and like PPV is derived from those with and without the condition of interest. The use of both PPV and NPV in research is cautionary since both are notably influenced by prevalence of the condition and may yield findings that are not clinically useful. As such, positive and negative likelihood ratios are generally recommended when determining post-test probability.

A positive likelihood ratio (LR+) is derived from subjects with and without the condition of interest. LR+ is calculated by taking the sensitivity and dividing by 1-specificity. With LR+, positive findings influence the post-test decision making and stronger LR+ will be notably greater than 1.0. In contrast, negative likelihood ratio (LR-) is calculated by taking 1-Sensitivity divided by the specificity, where a robust finding is hallmarked by smaller values closer to 0, and reflects a negative test finding only. Both LR- and LR+ are used to calculate post-test probabilities (0 to 100%) when the tests are negative or positive. A strong clinical test (or cluster of
tests) should have the ability to rule out a condition when negative (with post-test probabilities near 0) or rule in a condition when positive (with post-test probabilities near 100).

Further, truly robust tests should have confidence intervals that are precise, which is suggestive that repeating the study findings should lead to similar results. A confidence interval is a parameter estimate that outlines the boundaries that a given test value will fall if the study is repeated numerous times. In most cases, 95% confidence intervals are reported. This means that there is a 95% chance that the test value would fall within the boundaries of the confidence interval if the study was replicated in a different sample. For example, if a LR+ was 1.5 and the 95% confidence interval reported was 1.2 to 1.8, this means that there is a 95% chance that another trial would a LR+ between the boundaries of 1.2 to 1.8.

Because LR+ and LR- determine the values of both positive and negative findings, evaluate individuals with and without the condition of interest, and can be used to estimate post-test probabilities with adjustments for pre-test prevalence, these metrics should be used to guide decision making over sensitivity and specificity. Both LR+ and LR- facts on both those with and without the condition of interest and only these truly reflect a situation of diagnostic uncertainly.

**MULTIVARIATE MODELING**

The goal in any data analysis is to extract from raw information the accurate estimation (Alexopoulos, 2010). The goal when clustering tests is to determine the best combination estimates and to do so multivariate modeling is required. Thus, clustering is simply the act of evaluating a set of tests and measures together, in combination, when
making a clinical decision or a mathematical assessment. For example, Wainner and colleagues realized that there is no single test or measure that is strong enough to diagnosis cervical radiculopathy; however, when tests such as the upper limb tension test, Spurling’s test, cervical distraction, and assessment of range of motion to the affected side are clustered, the diagnostic accuracy is very strong. Clustering tests more closely reflects how many clinicians make decisions because it takes into account a number of presentations from the clinical assessment.

Multivariate modeling is a form of statistical analysis that explores the relationship between two or more predictor variables (the clinical tests) and the outcome variable (the reference standard). There are multiple forms of multivariate modeling methods and for clustering best tests and measures for diagnosis, a logistic regression analysis is the most appropriate type since the diagnosis is almost always dichotomous (present or absent).

Following proper multivariate modeling methodology is essential and the failure to do so when developing clustered rules or guides has been recognized by many authors (Beattie & Nelson, 2006; Beneciuk, Bishop, & George, 2009; C. Cook, Shah, & Pietrobon, 2008; Haskins, Rivett, & Osmotherly, 2012; May & Rosedale, 2009; Nee & Coppieters, 2011; Stanton, Hancock, Maher, & Koes, 2010). Although multivariate modeling can be notably complex, before considering clustering tests to determine most parsimonious values it is useful to contemplate the following four principles: 1) determination of observations per variable, 2) linearity continuous measures, 3) assessment of conditional dependence of the predictor variables (also recognized as Variance Inflation Factor or Tolerance) and 4) appropriate stepwise modeling. The
following paragraphs will provide recommendations for each of these principles.

**Determination of Observations per Variable:** There are a number of ways to determine the appropriate observations per variable, or, in determining how many tests should be included as independent variables in the multivariate model. For the sake of clarity, an observation would be an individual who is enrolled in the sample for the diagnostic accuracy study. For simple univariate multinomial or logistic regression, Hosmer and Lemeshow (Hosmer & Lemeshow, 2000) have recommended a minimum observation-to-variable ratio of 10, but cautioned that a number this low will likely overfit (overly burden) a model. That said, it is wisest to adopt the preferred observation-to-variable ratio of 20 to 1 for stepwise, multivariate modeling, with full recognition that a ratio of 50 to 1 is likely to provide more precise results. Thus, if one wanted to include 4 tests in a cluster multivariate model, a total of 80 to 200 patients are recommended.

**Linearity of Continuous Measures:** In most cases, clinical tests of the shoulder are either “positive” or “negative”. In some cases, a positive finding is determined after a threshold score is ascertained from a continuous set of measures (e.g., a threshold of a particular range of motion score). When underlying tests have a continuous value, the linearity of that value must be evaluated prior to determining a threshold. Linearity is generally analyzed by plotting to identify potential curvilinear relationships. An example of a lack of linearity is the estimation of the relationship of one’s flexibility to injury. It is suggested that those who are overly inflexible and those who are excessively flexible, are more predisposed to an overuse injury that one who is in the middle ranges. If we were to evaluate the influence of flexibility toward injury on values such as these, values that are not linear, no significant relationship would occur. There are adjustments one can make if
a variable lacks linearity. One can create categories and enter the variable as ordinal data with a set of indicators (dummies). Or, one can modify the definition of a positive test to reflect the variability within the underlying data. Using our previous flexibility example, one could score inflexible and excessively flexible as ‘positive’ and those in the middle categories as ‘negative’.

**Assessment of Conditional Dependence of the Predictor Variables:** One possible reason why past studies have failed to outline clusters of findings is the concept of conditional dependence. Conditional dependence (Menten, Boelaert, & Lesaffre, 2008) occurs when a subsequent test finding is not dissimilar to the first test finding or when a series of tests actually measure the same thing and are positive together in clusters or negative together in clusters. During multivariate modeling, this dependence is routinely referred to as assessment of multicollinearity. One can assess multicollinearity through use of correlation matrixes, variance inflation factors (VIF) and tolerance values. A correlational finding of \( r > 0.7 \) between test variables can be used to assess the potential of multicollinearity. (Shen & Gao, 2008) A mean VIF close to 1 represents little collinearity, whereas 10 or greater is very poor and reflects very high collinearity. (Kutner, Nachtsheim, & Neter, 2004) Tolerance is the reciprocal of VIF thus values close to 0 are considered to have high collinearity. (Firth, 1993) If tests are conditionally dependent and are included in the multivariate model, there is a risk that the test will be removed from the final model and/or the test will remain in the model but will adversely influence the beta scores of the variables within the model. In layman’s terms, a beta score allows for a consistent and meaningful measure across different units for the relationship of an independent variable to the dependent variable. A notable example of conditional
dependence was the recent publication on clustered tests for cervical myelopathy by Cook and colleagues (C. Cook, et al., 2010). In the study, various forms of reflex testing were nearly always hyper-responsive at different areas (quadiceps, Achilles, brachioradialis, etc), but only one of the tests (positive brachioradialis) was included in the clustered model. Adding all the findings to the final model would not improve the accuracy and could alter the beta estimates.

*Appropriate Stepwise Modeling:* Stepwise modeling is not without controversy. (Wlikinson & Dallal, 1981) Stepwise regression modeling is an automatic procedure in which the choice of predictive variables is carried out until the strongest, most refined explanatory model is determined. Most commonly, univariate analyses for each single test to the reference standard is calculated. Univariate analyses with p values of <0.15 are generally included in a multivariate model since the tests interactions with other tests may yield diagnostic findings in the final analyses. Using the automated stepwise processes, most statistical software programs will calculate a single cluster of independent variables (tests) that are responsible for the best explanation. By analyzing the best number of positive findings (e.g., 1 of 4, 2 of 4, 3 of 4, etc) one can further determine the desired sensitivity and specificity of their created cluster.

**BEST TEST CLUSTERS**

Before presenting the best published test clusters, “best” needs to be put in context. “Best” as used in this manuscript, is defined as those combinations of tests with the strongest likelihood ratios from research with the highest quality. The quality of the tests clusters is judged by using the Quality Assessment of Diagnostic Accuracy Studies (QUADAS) document and using a 0-14 (0=lowest quality) scale (Whiting, et al., 2004). From our past experience (C. E. Cook & Hegedus, 2011; Hegedus, et al., 2008), those
studies scoring below 10/14 are full of design faults that make the likelihood ratios from those studies questionable and probably not repeatable in populations outside of those examined in the study.

Only 4 articles (Farber, Castillo, Clough, Bahk, & McFarland, 2006; Guanche & Jones, 2003; Litaker, Pioro, El Bilbeisi, & Brems, 2000; Park, Yokota, Gill, El Rassi, & McFarland, 2005) met our quality criteria and these articles reported on just 6 current clusters. The best test clusters currently available are summarized in Table 1.

Unfortunately, even these high quality studies have failed in some respect with regard to sample size, stepwise regression, conditional dependence, and linearity of continuous measures (Table 2). Closer examination of these 4 studies reveals some other interesting findings. With regard to rotator cuff tears, of note is that both test clusters (Litaker, et al., 2000; Park, et al., 2005) incorporate older age as a component and that the most diagnostic cluster (Park, et al., 2005) uses 2 tests, painful arc (Bak, et al., 2010; Litaker, et al., 2000; Michener, Walsworth, Doukas, & Murphy, 2009) and drop arm (Bak, et al., 2010), that have low specificity or sensitivity values resulting in likelihood ratios that approach 1.0. In addition, another set of the tests from this same study (Park, et al., 2005), infraspinatus and painful arc, are part of the diagnostic cluster for impingement, likely leading to diagnostic confusion between early stages of impingement and rotator cuff tears (latter stage of impingement). The 2 diagnostic clusters for labral tears, a difficult clinical diagnosis, come from a single study (Guanche & Jones, 2003) and are only moderately diagnostic when positive with positive likelihood ratios ranging between 2.67 and 5.43. Further, one of the clusters for labral tears incorporates the active compression test, a test of dubious value (Ebinger, Magosch,
Lichtenberg, & Habermeyer, 2008; McFarland, Kim, & Savino, 2002; Morgan, Burkhart, Palmeri, & Gillespie, 1998; Oh, Kim, Kim, Gong, & Lee, 2008; Walsworth, Doukas, Murphy, Mielcarek, & Michener, 2008). Finally, for examining anterior instability, the results of the study (Farber, et al., 2006) are likely influenced by the fact that the instability group was younger and was more likely to have a history of trauma. It is important to note that in the study by Farber et al., (Farber, et al., 2006) apprehension was used as a positive test and not reproduction of pain.

Despite the limitations of current literature on the diagnostic accuracy of test clusters to diagnose shoulder pathology, we thought it would be helpful to illustrate the best combinations through a case study (Figure).

Case Study

Our fictitious patient is 67 years old and has complained of shoulder pain of 4 months in duration. He reported that his pain initiated while walking his dogs (when they jerked the leash he held) but notes that the pain has progressed markedly over the last two months. He is able to raise his arm above his head (with pain) but has noted that his arm now aches consistently, with a more noticeable ache at night. Frequent use of ibuprofen helps modulate his pain but the effects are only temporary at best.

As a clinician, one might consider several possibilities, especially with the individual’s age, consistent pain, and traumatic onset. Tests with low LR- help “rule out” competing conditions thus one might choose to consider tests or clusters of tests for shoulder labrum tears, impingement, or a rotator cuff tear. As a reminder, clustering tests often leads to higher LR+ with a sacrifice of LR-, unless the cluster was mathematically designed as a screen.
Since we have 3 competing diagnoses, attempting to rule out one or two conditions would be prudent. Currently, there are no high quality clusters of screening tests for the labrum that would rule out this condition. In the absence of a labral tear test cluster with a LR- near zero, the clinician has 2 choices: 1. consider single test results with a LR- near zero or 2. attempt to rule in the condition with a cluster of test findings that has a high LR+. Only 1 OST, the biceps load II, comes from a high quality study and has a LR- near zero(Kim, Ha, Ahn, & Choi, 2001). Unfortunately, a second high quality study(Oh, et al., 2008) showed the test to have no ability to rule out (or in) a labral tear. Guanche et al(Guanche & Jones, 2003) reported a test cluster with a LR+ of 5.43 but this likelihood ratio is of only moderate assistance in diagnosing a labral tear which does not have an established set of signs and symptoms(Luime, et al., 2004) and a likely low prevalence, somewhere around 6%(Snyder, Banas, & Karzel, 1995). Therefore, the best decision in this case is to treat the diagnosis of labral tear as one of exclusion and move on to rule out one of the diagnoses of either rotator cuff tear or impingement.

Park and colleagues(Park, et al., 2005) reported that with impingement syndrome, negative findings of 1) Hawkins-Kennedy, 2) painful arc test and 3) infraspinatus test provides a very low LR- (0.17) and thus, has the capacity to rule out the condition. Park et al(Park, et al., 2005) also reported that ruling out a rotator cuff tear is possible with negative findings on 1) Age ≥ 60 2) painful arc test, 3) drop arm test, and 4) infraspinatus test. Since our patient is over age 60, we cannot rule out a rotator cuff tear. For the sake of this case, the Hawkins-Kennedy and the painful arc tests were negative but the infraspinatus test (weakness against resisted external rotation) was positive. The impingement test cluster, therefore, cannot rule out
impingement. With a negative Hawkins-Kennedy and painful arc, we also cannot rule in impingement.

To rule in a rotator cuff tear, one could refer back to the findings of Park and colleagues (Park, et al., 2005) or consider the results from Litaker et al (Litaker, et al., 2000). Two consistencies with the findings of Park et al (Park, et al., 2005) and Litaker et al (Litaker, et al., 2000) are age and external rotation strength losses. Park et al (Park, et al., 2005) report the benefit of the drop arm sign and the painful arc test whereas Litaker et al (Litaker, et al., 2000) report the value of night pain. Recall that our patient is 67 years old has night pain, and a positive infraspinatus test (weakness in external rotation). These 3 findings complete the test cluster by Litaker et al (Litaker, et al., 2000). The clinician must be content with a LR+ of 9.84. Since the painful arc test was previously reported as negative, the cluster of Park et al (Park, et al., 2005), with a LR+ of 28.0, cannot be used.

**CONCLUSION**

The clinical diagnostic process should be viewed through the lens of odds and probabilities. In order to do so, test clusters from high quality studies should be utilized. In our case study, the patient likely has a rotator cuff tear but we were unable to rule out or in a labral tear and impingement. High quality clinical test clusters with powerful diagnostic characteristics for labral tears do not presently exist and impingement is an all-encompassing term for tendon pathology at the shoulder that is, at best, unhelpful in guiding treatment, and, at worst, a clinical illusion (Hegedus, et al., 2012; Lewis, 2011). Other important pathologies of the shoulder like biceps tendinopathy, multi-directional instability, and fractures, also lack powerful clinical diagnostic clusters. Improved research that follows the tenets
of multivariate modeling outlined in this article must be performed in order to improve the tools available to clinicians as we attempt to make use of the clinical examination to diagnose shoulder pathology.

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Ethical Approval: N/A

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Table 1. Best test clusters from current literature

<table>
<thead>
<tr>
<th>Author(s)</th>
<th>Pathology</th>
<th>Test Cluster</th>
<th>LR+</th>
<th>LR-</th>
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</thead>
<tbody>
<tr>
<td>(Litaker, et al., 2000)</td>
<td>Rotator cuff tear</td>
<td>1. Age &gt; 65 and 2. Weakness in external rotation and 3. Night pain</td>
<td>9.84</td>
<td>0.54</td>
</tr>
<tr>
<td>(Park, et al., 2005)</td>
<td>Rotator cuff tear (full thickness)</td>
<td>1. Age ≥ 60 and 2. + painful arc test and 3. + drop arm test and 4. + infraspinatus test</td>
<td>28.0</td>
<td>0.09</td>
</tr>
<tr>
<td>(Park, et al., 2005)</td>
<td>Impingement</td>
<td>1. + Hawkins-Kennedy and 2. + painful arc test and 3. + infraspinatus test</td>
<td>10.56</td>
<td>0.17</td>
</tr>
<tr>
<td>(Farber, et al., 2006)</td>
<td>Anterior instability (traumatic)</td>
<td>1. + apprehension test and 2. + relocation test</td>
<td>39.68</td>
<td>0.19</td>
</tr>
<tr>
<td>(Guanche &amp; Jones, 2003)</td>
<td>Labral tear</td>
<td>1. + relocation test and 2. + active compression test</td>
<td>4.56</td>
<td>0.65</td>
</tr>
<tr>
<td>(Guanche &amp; Jones, 2003)</td>
<td>Labral tear</td>
<td>1. + relocation test and 2. + apprehension test</td>
<td>5.43</td>
<td>0.67</td>
</tr>
</tbody>
</table>

Table 2. Design features of the best articles reporting on diagnostic accuracy of combined tests.

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<tbody>
<tr>
<td>At least 20 subjects per test in the cluster</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
<td>yes</td>
</tr>
<tr>
<td>Assessed conditional dependence</td>
<td>yes</td>
<td>no</td>
<td>no</td>
<td>no</td>
</tr>
<tr>
<td>Stepwise regression</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
<td>no</td>
</tr>
<tr>
<td>Assessed linearity of continuous measures</td>
<td>yes</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
</tbody>
</table>
Figure. Diagnostic process using the best available clinical test clusters for shoulder pathology

Key: + = positive test; - = negative test; H-K = Hawkins-Kennedy; ER = external rotation

Based on history, 3 competing diagnoses:

Rule out: Labral tear - NO

Rotator cuff tear

Impingement

Labral tear

Rotator cuff tear - NO (age)

Impingement - NO
(+ infraspinatus test)

Rule in: Labral tear - NO

Rotator cuff tear - YES!
1. Age > 65
2. ER weak
3. Night pain

Impingement - NO
(-H-K & painful arc)

Working diagnosis: Rotator cuff tear


