Health Technology Clinical Committee  
**Date:** November 14, 2008  
**Time:** 8:00 am – 5:00 pm  
**Location:** Marriott Hotel – 3201 South 176th Street, Seattle, WA 98188  
**Teleconference Bridge:** 1-360-923-2997  
**Access Code:** 360-946-1464

*D*R*A*F*T*  
HTCC MINUTES

**Members Present:** Brian Budenholzer; Michael Myint; Carson Odegard; Richard Phillips; Michelle Simon; Lydia Bartholomew; Jay Klarnet; C. Craige Blackmore; Michael Souter and Louise Kaplan.

**HTCC FORMAL ACTION**

1. **Call to Order:** Dr. Budenholzer, Chair, called the meeting to order at 8:00 a.m. Sufficient members were present to constitute a quorum.

2. **Implantable Infusion Pump Findings and Decision:** Dr. Budenholzer referred members to the draft findings and decision and called for further discussion or objection. The committee agreed to insert a statement under the Committee Decision portion of the Findings and Decision that states: “the committee has received the Implantable Infusion Pump public comments through October 10th at the October 17th and November 14th, 2008 public meeting and have incorporated those comments in finalizing their decision.”
   
   ➢ **Action:** Six committee members unanimously approved the Implantable Infusion Pump findings and decision document. Two committee members abstained from voting. One committee member wasn’t present at the time the vote was conducted.

   ▪ HTA program staff will edit any typos before finalizing.

3. **Computed Tomographic Angiography Determination:** The HTCC reviewed and considered the Computed Tomographic Angiography for detection of coronary artery disease technology assessment report, information provided by the Administrator, state agencies, and public members; and heard comments from the evidence reviewer, HTA program, agency medical directors, and several public members. The committee considered all the evidence and gave greatest weight to the evidence it determined, based on objective factors, to be the most valid and reliable.
HTCC COMMITTEE COVERAGE DETERMINATION VOTE

<table>
<thead>
<tr>
<th></th>
<th>Not covered</th>
<th>Covered Unconditionally</th>
<th>Covered Under Certain Conditions</th>
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</thead>
<tbody>
<tr>
<td>Computed Tomographic Angiography</td>
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<td>0</td>
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- Conditions for coverage: Investigation of acute chest pain in an emergency department or hospital setting who are at low-to-intermediate risk of coronary artery disease. Type of technology to be used is a 64-slice or better. The committee unanimously agreed on these conditions.

- Action: The committee chair directed HTA staff to prepare a Findings and Decision document on Coronary Computed Tomographic Angiography reflective of the majority vote for final approval at the next public meeting.
SUMMARY OF HTCC MEETING TOPICS, PRESENTATION, AND DISCUSSION

Agenda Item: Welcome & Introductions

✓ The Health Technology Clinical Committee (HTCC) met on November 14, 2008.

Agenda Item: Meeting Open and HTA Program Update

Dr. Brian Budenholzer, HTCC Chair opened the public meeting. Leah Hole-Curry, HTA Program Director, provided an overview of the agenda, meeting guide and purpose, room logistics, and introductions.

Leah Hole-Curry, HTA Program Director, provided an update on HTA program activities and outcomes.

✓ 2009 - Potential Topics will be referred to the Administrator for his consideration, technologies include: Glucose Monitoring, Sleep Apnea Diagnosis and treatment, Calcium Scoring for cardiac disease, Vagal Nerve Stimulation, Elective Cesarean Section, Hip Resurfacing, Osteoarticular Transfer System – Cartilage Surgery (OATS procedure), Bone Growth Stimulators, Massage Therapy for Chronic Head, Neck and Back pain, Transcutaneous Electrical Neural Stimulation (TENS procedure), Essure Permanent Birth Control procedure, and Breast Cancer Tumor Screening.

✓ Clinical Committee Recruitment: The HTA program announced that we have an available recruitment on the health technology clinical committee. Daniel Abrahamson resigned from his clinical committee membership at the October 17, 2008 public meeting. The program is working towards filling this open recruitment.

✓ Artifical Disc Replacement: the HTA program is in the process of drafting a Findings and Decision on Artificial Disc Replacement. The program will publish post the Findings and Decision to the web and will circulate to the committee members as for comments. The health technology clinical committee will adopt the Findings and Decision at their next public meeting.

Agenda Item: Previous Meeting Business

✓ Overview of the draft minutes from the October 17, 2008 public meeting - the minutes are in the process of being drafted by HTA staff, they will then be posted to the web and circulated to committee members for comments. The health technology clinical committee will adopt the October 17th minutes at their next public meeting.

✓ Implantable Infusion Pumps Findings and Decision: Dr. Budenholzer referred members to the draft findings and decision and called for further discussion or objection. The committee agreed to insert a statement under the Committee Decision portion of the Findings and Decision that states: “the committee has received the Implantable Infusion Pump public comments through October 10th at the October 17th and November 14th, 2008 public meeting and have incorporated those comments in finalizing their decision.”

✓ Committee members commented on draft decision and public comments received:
  ▪ The public comments do not address or challenge the evidence the committee relied upon and the technology report
  ▪ Still has concerns regarding safety issues and discrimination between cancer/non-cancer patients. Public comments reinforced personal view, but majority vote is reasonable
the public comments were not based primarily on evidence; committee made a reasonable decision. Chronic pain vs. cancer there is a difference in timing and diagnosis is fatal, and the use for cancer was not before the committee. Comment from widow reinforced safety issues.

- Updated draft decision clarified and laid out decision well. For emerging trials, can re-review if necessary.
- Empathetic emotionally to public comments related to alleviation of pain, but committee made a rational decision based on evidence from a critically appraised report
- Disagree with public comments that statutory burden for committee decision is not met; updated findings reflect rationale and evidence.
- Overriding concern about serious adverse events occurring; case series efficacy evidence does not prove it to be effective are very low quality. Guidelines subsequently provided do not cite new reliable evidence; most cited studies were analyzed and rejected by ECRI;
- An open and transparent process was used, technology assessment included complete and relevant information;
  - **Action:** Six committee members unanimously approved the Implantable Infusion Pump findings and decision document. Two committee members abstained from voting.

**Action:** HTA program staff will edit any typos before finalizing.

**Agenda Item: Computed Tomographic Angiography Topic Review**

Leah Hole-Curry, HTA Program Director, introduced the primary technology topic to discuss:

- **Computed Tomographic Angiography for detection of coronary artery disease:** review of the evidence of the safety, efficacy and cost-effectiveness of Computed Tomographic Angiography.

**Computed Tomographic Angiography**

- Heart disease is the leading cause of death and disability in US: with 700,000 deaths. The most common heart disease in the US is coronary artery disease (CAD), which can lead to heart attack. CAD is a narrowing of one or more coronary arteries that result in an insufficient supply of oxygen to the heart muscle and is a leading cause of death in the US and developed countries.
  - CAD may be asymptomatic or lead to chest pain (angina), heart attack, myocardial infarction (MI) or death.
- Cardiac related diagnostic tests include both non-invasive and invasive tests.
  - Non invasive tests include: Stress Echocardiograms – tests that compare blood flow with and without exercise and visualize the heart. Single-photon emission computed tomography (SPECT), also known as nuclear stress testing or myocardial perfusion imaging.
  - Invasive tests include: The “gold” standard is the conventional coronary angiography which involves placement of a catheter and injection of contract material into a large artery or vein, followed by 2-dimensional visualization with x-rays.
- CCTA involves the use of CT scans and an injected dye to develop computer-aided, 3-dimensional images of the artery.
CCTA Potential Benefits: multiple-angle and multiple-plane visualization; improved visualization of soft tissues and adjacent anatomy; and lower degree of invasiveness compared to conventional CA.

CCTA Potential Drawbacks: increased radiation exposure; the possibility of incidental findings in adjacent anatomic structures; and the need for further testing (additive rather than replacement test).

CMS Decisions and Expert Treatment Guidelines

- Centers for Medicare and Medicaid Services (2008): no national coverage decision (NCD). Coverage memo conclusions: in summary, there is uncertainty regarding any potential health benefits or patient management alterations from including coronary CTA in the diagnostic workup of patients who may have CAD. No adequately powered study has established that improved health outcomes can be casually attributed to coronary CTA for any well-defined clinical indication, and the body of evidence is of overall limited quality and limited applicability to Medicare patients with typical co-morbidities in community practice. The primary safety concerns with cardiac CTA are the exposure to radiation and the use of contrast and blocker medications.

- American Heart Association (2006): evidence supports the use of CCTA for patients with low-to-intermediate stenosis and may obviate the need for ICA.

- Multi-Society Statement of Appropriateness Criteria for Cardiac Computed Tomography (2006): Appropriateness reviews deemed the use of CCTA for detection of CAD to be appropriate for the following patient populations: chest pain syndrome with intermediate pre-test probability of CAD and uninterpretable EKG or inability to exercise; chest pain and uninterpretable or equivocal stress test results; acute chest pain with intermediate pre-test probability of CAD and no EKG changes and serial enzymes negative; and symptomatic patients requiring evaluation of suspected coronary anomalies.

- American College of Radiology (2006): CCTA is appropriate for assessment of CAD, although its usefulness for patients with low pre-test probability is unknown. Appropriateness rating of 7 out 9 for the evaluation of chronic chest pain.

- SCCT/NASCI Consensus Update (2007): CCTA to be appropriate in the following circumstances: (1) to rule out significant coronary stenosis; (2) to evaluate patients with equivocal or discordant results on a stress perfusion or wall motion study; (3) to rule out stenosis in patients with a low pre-test likelihood of CAD and (4) to potentially replace diagnostic catheterization in patients undergoing non-coronary cardiac surgery.

**Agenda Item: Public Comments**

- Scheduled Public Comments: No scheduled public comments.

- Open Public Comments: Four individuals provided comments during the open portion (limited to three minute comments) –
  - Dr. Kelley Branch (University of Washington); Dr. William Shuman (University of Washington); and Dr. Edham Ward: provided a statement approving the use of CCTA.

**Agenda Item: Computed Tomographic Angiography Topic – Agency Data**
Dr. Malcolm Dejnozka, Uniform Medical Plan Medical Director, presented to the committee the agency utilization and outcomes for Computed Tomographic Angiography.

- **Key agency concerns for prioritization:**
  - Efficacy concerns – High: evidence of CCTA sensitivity, specificity, reliability are mixed. Rapid technology evolution/diffusion; what’s the community standard outside the research or “center” experience; interpretation reliability (inter-rater reliability) concerns; and patient selection. Gold standard exists = invasive coronary angiography.
  - Safety concerns – Low: Short-term ~ IV contrast reaction; renal insufficiency; procedure drugs (beta-blockers/nitrates) and dilemma of “incidental findings” which may add potentially harmful tests / procedures. Long-term ~ radiation exposure is significant; especially if a screening tool.
  - Cost concerns – High: economic impact of CAD is greater than $120 billion (2004). Does CCTA add costs, drive other costs or eliminate need for alternative tests? Half-life of new generation CT propagates market change (variation) and costs.

- **Agency coverage experience (CCTA) – coverage policies vary by agency:**
  - L&I: not within scope of services.
  - UMP: deemed “investigational” for most uses, but considered by exception by pre-authorization and medical review.
  - DSHS: covered, requires pre-authorization.
  - The agencies cover alternatives (coverage policies vary by agency): CABG; SPEC (i.e., nuclear medicine stress test); STRESS ECHO and Invasive Coronary Angiography.

- **Data query to CY 2006 and 2007 – claims DATA base query CPT code constraints include:**
  - Patients may get one, or multiple studies; mixed primary and secondary payer (to Medicare) costs; network / non-network rates; health plan participants (changes in demographics); and site of service.

- **Washington State Agencies Experience:**

  **Invasive Coronary Angiography**

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PEHP*: Costs are skewed: UMP was secondary for 1,350 patients covered under Medicare; 1,405 patients UMP primary (average $1,893.99 paid).

**STRESS ECHO Utilization**

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*PEHP* Costs are skewed: UMP was secondary for 7,665 patients covered under Medicare; 5,930 patients UMP primary (average $265.09 paid).

## Coronary CT Angiography Utilization

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*PEHP* Costs are skewed: UMP was secondary for 4,920 patients covered under Medicare; 6,541 patients UMP primary (average $614.73 paid).

## Summary of Overall Costs

<table>
<thead>
<tr>
<th>Agency</th>
<th>Patients</th>
<th>ICA</th>
<th>Stress Echo</th>
<th>SPECT</th>
<th>CCTA</th>
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## Coronary CT Angiography – ICER and Agency Utilization

<table>
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<tr>
<th>Procedure</th>
<th>ICER estimates Total ED Costs (Relative Ratio)</th>
<th>Average Agency Costs (Relative Ratio)*</th>
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<tr>
<td>SECHO</td>
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<tr>
<td>ICA</td>
<td>$2,750 (5.90)</td>
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<tr>
<td>SPECT</td>
<td>$765 (1.64)</td>
<td>$409 (0.59)</td>
</tr>
<tr>
<td>CCTA</td>
<td>$466 (1.00)</td>
<td>$687 (1.00)</td>
</tr>
</tbody>
</table>

ICER: “Threshold CCTA cost for cost savings in ED = $762”
Agency Conclusions: Cardiac Imaging for CAD is extensive – imaging options are available and competitive; technology use rapidly disseminating and evolving (“snapshot”); and screening not TEC assessed. Safety and potential harms – less invasive, but subjects’ patients to radiation exposure (long-term cancer risk) and dilemma of incidental findings (added studies/interventions). Costs analysis – moderate stenosis reassuring to clinicians/patients or generate an “oculostenotic reflex” (i.e., aggressive tests/treatments); cost advantage seen in the ICER systematic report might be offset by real-world reimbursements and “incidental findings” tests; and cost analytical model shouldn’t be generalized outside ED Triage setting.

- Evidence is most supportive in the ED Triage care setting. Insufficient evidence in other settings.

Agenda Item: Evidence Review Presentation

ICER presented an overview of their evidence report.

- Scope: CCTA technology 64-slice or better precision, reports between 2005 and present evaluated. CCTA use in emergency department triage of acute chest pain and outpatient evaluation of patients with stable chest pain and low-to-intermediate CAD risk.

- Coronary computed tomographic angiography (CCTA) is a minimally invasive radiological technique used to provide images of the heart and surrounding vessels. CCTA has been suggested as an alternative or useful complementary approach to other non-invasive methods of diagnosing coronary artery disease (CAD). Due to its ability to visualize coronary anatomy, CCTA has been suggested as a strategy to rule out significant CAD among patients at low or intermediate risk of significant disease, thereby giving greater reassurance than other non-invasive methods and potentially reducing the number of patients ultimately sent for invasive coronary angiography (ICA).

  - ICA is typically an inpatient procedure. At the time of the procedure a catheter is inserted into an artery, usually the femoral blood vessel, and contrast dye is injected through the catheter. X-ray images are then captured and displayed on a video screen (a procedure known as fluoroscopy), and can be viewed either as images or in motion picture form. While complications from ICA are relatively infrequent, they can be significant, and include myocardial infarction, cardiac arrhythmia, stroke, hemorrhage, infection, trauma to the artery from hematoma or from the catheter, sudden hypotension, and reaction to the contrast medium.

  - Stress echocardiograms (ECHO) produce images of the heart through the use of sound waves. The test allows for the evaluation of blood flow in different areas of the heart to identify weak or damaged areas of the muscle. This is done through a comparison of images at rest and under cardiac stress induced by exercise or pharmacologic means. Clinically, the test is simple to perform, relatively inexpensive, and easily accessible. However, the image quality is lower in obese patients and those with chronic disease.

  - SPECT imaging involves the use of a tracer radiopharmaceutical to highlight areas of decreased blood flow in the myocardium. Images are captured via a gamma camera, and...
may be reconstructed to create two or three-dimensional films. The accuracy of SPECT imaging has improved to the point that it is often used for prognostic use in addition to diagnosis. SPECT also involves the use of contrast media and delivers a radiation dose similar in magnitude to that of ICA and CCTA.

**CCTA** is a technique in which a CT scanner is used to acquire multiple simultaneous tomographic sections (“slices”) of the coronary arteries. At the time of this outpatient procedure, an IV is placed into a peripheral vein and a contrast dye is administered for the purposes of visually defining the arteries for the scan. Beta blockers may be given to the patient to slow the heart rate in order to prevent artifacts of heart motion that may affect image quality. The patient is positioned on the CT scanner and a large number of x-ray images are taken from multiple angles and reconstructed using computer software. Multi-detector row CT scanners contain rotating gantries that capture multiple images, or “slices”. A 64-slice CCTA was introduced in 2004 and increased the number of captured images from the previous 16- and 32-slice technology. The 64-slice scanner has rapidly replaced earlier versions and is currently considered to be the community standard for CCTA.

- In the emergency department, CCTA can be used for the triage of patients experiencing acute chest pain to “rule out” CAD as the underlying cause.
- In the outpatient setting, CCTA is most often used to evaluate patients with stable, non-emergent symptoms. For such patients CCTA can be used as an initial test or as a method for further evaluation following inconclusive results from another non-invasive functional test.

**Compared to other non-invasive diagnostic methods** there are also potential disadvantages specific to CCTA, including a small risk of allergic reaction from the use of contrast dye and the risk of renal damage from the dye among patients with pre-existing renal dysfunction. In addition, the increased precision from multi-detector row CT scanners is accompanied by a higher radiation dose to the patient. Lastly, the range of visualization of CCTA extends beyond the heart itself, creating the possibility of identification of “incidental findings” that may or may not be related to the patients’ complaints of chest discomfort.

**Description of Included Studies:**

- **ED** – 8 studies met criteria (N=686); age range = 46 to 58 years; 1 RCT, others single-center case series; and most used clinical diagnosis algorithm for confirmation.
- **Outpatient** – 34 studies met criteria (N=3,349); age range = 46 to 69 years; and most used ICA alone or in combination as referent.

**Potential Harms – Radiation Exposure:** effective dose reported in 17 studies; overall range = 4.6 to 21.4 mSv; lowest rates reported for studies using dose-sparing protocols or dual-source scanners. Six studies reported separate doses for men and women – Men = 7.45 – 15.2 mSv (mean 12.4); Women = 10.24 – 21.4 mSv (mean 14.2).

**Evidence review - conclusion interpretation for what is known**

- For asymptomatic: no professional support or evidence for use as asymptomatic screening test
- For High risk patients: professional guidelines support high-risk patients going directly to invasive catheterization rather than through this test
For low-intermediate risk patients in the ED - Diagnostic accuracy of 64-slice as triage tool supported by one RCT and several case series. Modeling suggests that under most assumptions CCTA is cost-saving.

For low-intermediate risk outpatients - No RCT evidence, no long-term cohort evidence Diagnostic accuracy of 64-slice appears very good compared to ICA and better at identifying occlusion than other non-invasive tests.

Modeling suggests lower rate of false positives than SECHO and SPECT, and lower rate of false positives than SPECT, but differences change with underlying prevalence of CAD and involves other trade-offs.

Conclusion interpretation: What we don’t know from the evidence

- Does CCTA change clinician threshold for testing?
- Does CCTA change physician decision-making in the outpatient setting?
- Does CCTA reduce anxiety or repeat testing?
- Does CCTA reduce invasive catheterization rates?
- Are incidental findings a benefit or harm?
- What is the impact of radiation exposure?
- Does treatment of CAD identified by CCTA among low-risk populations bring same benefits as treatment of CAD in prior studies?

**Agenda Item:** HTCC Computed Tomographic Angiography Discussion

Brian Budenholzer, Committee Chair, led a discussion of the evidence related to the safety, efficacy, and cost effectiveness of Computed Tomographic Angiography beginning with identification of key factors and health outcomes, and then a discussion of what evidence existed on those factors.

**Evidence availability and technology features**

The committee finds the following key factors relevant to the coverage decision:

- The evidence based technology assessment report indicates that Coronary Artery Disease (CAD) is burdensome and costly; an important and common medical concern. Detecting CAD and appropriately identifying the level of disease is important to select an appropriate treatment and prevent chest pain, heart attack and stroke. There are established non-invasive and invasive tests for detection of CAD. The gold standard comparison is to invasive coronary angiography (ICA); and non-invasive comparators include Stress ECHO and SPECT. The potential benefit of reducing the use of ICA is that it can reduce risks associated with an invasive procedure and reduce usage of associated percutaneous coronary treatment interventions. ICA mortality rates are 1 in 1000 and morbidity from stroke, infection, or bleeding are between .2 and .3%. Potential risks to using CCTA are that the test may be additive rather than a replacement which increases overall risks and costs; radiation exposure; unclear effect on health outcomes of patients and effect of incidental findings.

- The committee agreed with the technology assessment findings that there is not current evidence nor professional support for using CCTA in either asymptomatic populations (screening) or very high risk patients who are sent directly to catheterization. The committee further agreed with the
technology assessment distinction by separating discussion of use of CCTA into emergency department use and outpatient use, with a focus on the potential efficacy, safety and costs impacts in populations at low to intermediate risk of CAD.

The evidence based technology assessment report searched peer reviewed medical literature, submitted comments and other sources and identified eight trials, including one randomized controlled trial for emergency department use of CCTA; and 34 trials (no randomized) addressing outpatient usage of CCTA. Finally, the report included four cost studies addressing use of CCTA in ED and outpatient settings.

Key Factors and Health Outcomes Considered – Computed Tomographic Angiography

**Efficacy:** The committee identified multiple key factors and health outcomes that were important for consideration in their overall decision on whether the technology was effective. Summary of committee consideration, discussion, and comments are listed below.

- **Diagnostic Accuracy – Sensitivity:** the committee agreed as a whole that CCTA has a high level of sensitivity. The technology report sensitivity rate was 98%; which compared favorably to stress echo at 76-94% and SPECT at 88-98%. The indeterminate rates were also lower, with CCTA at 3% versus Stress ECHO at 13% and SPECT at 9%.

- **Diagnostic Accuracy – Specificity:** the committee agreed equivalent specificity. Some uncertainty about lower prevalence population was shared amongst the committee members. The technology report specificity rate was comparable at 82-88%; compared to stress echo at 88% and SPECT at 77%.

- **Reduction in invasive CA:** the committee agreed that modeling suggests reduced ICA, but trial evidence data was inconclusive with Rubenstien trial showing reduction and Goldstein shiwioing slight increase, especially when compared to alternative diagnostic tools.

- **Replace other tests:** most modeled analysis and clinical trials used CCTA in conjunction with other tests. Committee agreed that CCTA wouldn’t replace other non-invasive technologies.

- **Incidental findings:** committee discussed as an issue both we respect to efficacy and safety and concluded that evidence demonstrates incidental findings are not infrequent events. Incidental findings can provide valuable information for diagnosis of previously undetected other diseases but also often leads to uncertainty or further tests to rule out questionable findings. The committee agreed that there is currently no evidence regarding improved patient health outcomes balancing cost and potential harms from further testing and anxiety.

- **Effect in real world:** Committee discussed several technology assessment key unknowns: whether more disease found will help or harm patients, especially at lower disease levels (clinical relevance is questionable); whether broad dissemination will result in lower test thresholds that may not result in better overall health outcomes but more radiation; and the extent to which CCTA can replace and not add to tests. Additionally, certification of machines and readers was also discussed, hospitals require JAHCO accreditation and thus have some certification requirements.

**Safety:** The committee identified multiple key factors and health outcomes that were important for consideration in their overall decision on whether the technology was safe. Summary of committee consideration, discussion, and comments are listed below.

- **Radiation Exposure** is an important safety outcome to the committee. The committee discussed the technology assessment report findings of an overall cancer risk of .22% for women and .08% for men. Radiation dosage can be reduced through technique and machine type, but unknown whether these lowest dosage techniques/machines are used in WA settings.
Overall exposure reported at between 2.0-8.0mSV for lower range is equivalent to SPECT; and 12.0 to 14.0 range for higher dose which is equivalent to A-bomb survivor at 2.3 kilometer distance. The committee concluded that there are small but finite risks, within appropriate norms. The radiation risks are high enough to obviate benefit when applied to very low risk patients.

- **Incidental findings:** committee discussed as an issue both we respect to efficacy and safety and concluded that evidence demonstrates incidental findings are not infrequent events. Incidental findings can provide valuable information for diagnosis of previously undetected other diseases but also often leads to uncertainty or further tests to rule out questionable findings. The committee agreed that there is currently no evidence regarding improved patient health outcomes balancing cost and potential harms from further testing and anxiety.

**Cost:** The committee discussed cost and cost-effectiveness as a whole. This topic generated the least discussion. There are several cost models for ED and outpatient showing cost savings. The technology assessment report also modeled costs for ED and outpatient showing cost savings using Medicare reimbursement rates. No analysis included costs related to incidental findings or harms. Current state agency reimbursement rates do not correlate with model costs (Agency reimbursement for CCTA is higher and for comparators is lower).

- Committee members were split, with four considering the cost effectiveness currently unproven and five concluding that CCTA is either equivalent or more cost effective in some situations.

**Medicare Decision and Expert guidelines**
Committee reviewed and discussed the Medicare coverage decision and expert guidelines as identified and reported in the technology assessment report.

- There is no national coverage decision (NCD), however a coverage analysis and memo was issued in 2008 and summarized: there is uncertainty regarding any potential health benefits or patient management alterations from including coronary CTA in the diagnostic workup of patients who may have CAD. No adequately powered study has established that improved health outcomes can be casually attributed to coronary CTA for any well-defined clinical indication, and the body of evidence is of overall limited quality and limited applicability to Medicare patients with typical co-morbidities in community practice. The primary safety concerns with cardiac CTA are the exposure to radiation and the use of contrast and blocker medications.

- Four expert guidelines were identified that address the use of CCTA for detection of CAD, but not the setting (ED versus outpatient).
  - American Heart Association (2006): evidence supports the use of CCTA for patients with low-to-intermediate stenosis and may obviate the need for ICA.
  - Multi-Society Statement of Appropriateness Criteria for Cardiac Computed Tomography (2006): Appropriateness reviews deemed the use of CCTA for detection of CAD to be appropriate for the following patient populations: chest pain syndrome with intermediate pre-test probability of CAD and uninterpretable EKG or inability to exercise; chest pain and uninterpretable or equivocal stress test results; acute chest pain with intermediate pre-test probability of CAD and no EKG changes and serial enzymes negative; and symptomatic patients requiring evaluation of suspected coronary anomalies.
  - American College of Radiology (2006): CCTA is appropriate for assessment of CAD, although its usefulness for patients with low pre-test probability is unknown. Appropriateness rating of 7 out 9 for the evaluation of chronic chest pain.
SCCT/NASCI Consensus Update (2007): CCTA to be appropriate in the following circumstances:
(1) to rule out significant coronary stenosis; (2) to evaluate patients with equivocal or discordant
results on a stress perfusion or wall motion study; (3) to rule out stenosis in patients with a low
pre-test likelihood of CAD and (4) to potentially replace diagnostic catheterization in patients
undergoing non-coronary cardiac surgery.

Agenda Item: Computed Tomographic Angiography Vote

The clinical committee utilized their decision tool to first gauge committee judgment on the status of
the evidence in the three primary areas of safety, efficacy, and cost.

Computed Tomographic Angiography Votes:

Is there sufficient evidence under some or all situations that the technology is:

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<tr>
<th></th>
<th>Unproven (no)</th>
<th>Equivalent (yes)</th>
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<tr>
<td>Effective</td>
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<td>6</td>
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<td>Safe</td>
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<td>Cost-effective</td>
<td>4</td>
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Committee Discussion related to ad hoc group. Committee discussed whether an ad hoc group
was needed to provide more information to the committee:
- Review of literature is well done; information is present to make decision; there is nothing out
  there that would likely change view.
- Ad hoc committee would provide more opinion, but committee has good idea of what
  information is available and what the evidence is.
- We have all the information before us; we just need to make a decision.

HTCC Computed Tomographic Angiography Decision

The HTCC reviewed and considered a comprehensive 2008 HTA Evidence Report on Computed
Tomographic Angiography that included and analyzed the relevant and highest quality studies. The
committee also reviewed information provided by the Administrator, state agencies, and public
members; and heard comments from the evidence reviewer, HTA program, agency medical director,
and several public members.

Based on the evidence provided and the information and comments presented, the committee moved to
a vote on coverage.

<table>
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<th>HTCC COMMITTEE COVERAGE DETERMINATION</th>
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<td></td>
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<tr>
<td>Computed Tomographic Angiography</td>
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<tr>
<td>Not covered</td>
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<tr>
<td>Covered Unconditionally</td>
</tr>
<tr>
<td>Covered Under Certain Conditions</td>
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Committee Discussion related to Expert Treatment Guidelines and Medicare Decision:

- There is no national Medicare coverage decision. The decision is consistent with treatment guidelines in that low to intermediate triage will be covered, although the coverage decision is more specific in identifying the place of service.
- The committee decision is based on all evidence, including public and agency comments and the comprehensive technology assessment report.

Outcome: The committee chair directed HTA staff to prepare a Findings and Decision document on Computed Tomographic Angiography reflective of the majority vote for final approval at the next public meeting.

- CCTA with Conditions: Investigation of acute chest pain in an emergency room or hospital setting who are at low-to-intermediate risk of CAD.
  - Type of technology to be used is a 64-slice or better.
Health Technology Assessment - HTA

Health Technology Clinical Committee
Findings and Coverage Decision
Topic: Artificial Disc Replacement
Meeting Date: October 17, 2008
Final Adoption: *D*R*A*F*T updated

Number and Coverage Topic
10172008 - Artificial Disc Replacement

HTCC Coverage Determination
Cervical and Lumbar Artificial Disc Replacement is a covered benefit only under criteria identified in the reimbursement determination.

HTCC Reimbursement Determination
❖ Limitations of Coverage:

Lumbar ADR
1) Patients must first complete a structured, intensive, multi-disciplinary program for management of pain, if covered by the agency;
2) Patients must be 60 years or under;
3) Patients must meet FDA approved indications for use and not have any contra-indications. FDA approval is device specific but includes:
   • Failure of at least six months of conservative treatment
   • Skeletally mature patient
   • Replacement of a single disc for degenerative disc disease at one level confirmed by patient history and imaging

Artificial Disc Replacement FDA general contra-indications:
   • Active systemic infection or infection localized to site of implantation
   • Allergy or sensitivity to implant materials
   • Certain bone and spine diseases (e.g. osteoporosis, spondylosis)

Cervical ADR
1) Patients must meet FDA approved indications for use and not have any contra-indications. FDA approval is device specific but includes:
   • Skeletally mature patient
Reconstruction of a disc following single level discectomy for intractable symptomatic cervical disc disease (radiculopathy or myelopathy) confirmed by patient findings and imaging.

Artificial Disc Replacement FDA general contra-indications:
- Active systemic infection or infection localized to site of implantation
- Allergy or sensitivity to implant materials
- Certain bone and spine diseases (e.g. severe spondylosis or marked cervical instability)

Non-Covered Indications
Non-FDA approved uses

Agency Contact Information

<table>
<thead>
<tr>
<th>Agency</th>
<th>Contact Phone Number</th>
</tr>
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<tbody>
<tr>
<td>Labor and Industries</td>
<td>1-800-547-8367</td>
</tr>
<tr>
<td>Uniform Medical Plan</td>
<td>1-800-762-6004</td>
</tr>
<tr>
<td>Health and Recovery Services Administration</td>
<td>1-800-562-3022</td>
</tr>
</tbody>
</table>

A. Health Technology Background

The Artificial Disc Replacement (ADR) topic was selected and published in August 2007 to undergo an evidence review process per RCW 70.14.100(1)(a). ADR is the complete removal of the damaged disc and implantation of an artificial disc. The intent is to treat the pain and disability believed to be caused by a degenerated disc by removing the diseased. Both L-ADR and C-ADR are intended to preserve motion at the involved spinal level and therefore decrease stresses on adjacent segment structures and the risk of adjacent segment disease.

The HCA Administrator contracted with an independent technology assessment center for a systematic evidence based technology assessment report of the technology’s safety, efficacy, and cost-effectiveness consistent with RCW 70.14.100(4). On August 27, 2008, the HTA posted a draft report, invited public comment, and posted a final report on September 19, 2008. The contractor reviewed publicly submitted information, and searched, summarized, and evaluated trials, articles, and other evidence about the topic. This comprehensive, public and peer reviewed, report is 230 pages, identified 176 potentially relevant articles, and a Medicare coverage decision. Based on pre-established criteria and clinical research methodology, the technology assessment center included the most relevant and best available evidence on the safety, effectiveness, and cost effectiveness of artificial disc replacement. The result is a critical appraisal of: 2 moderate quality randomized controlled trials, 25 low quality case series, and 2 economic analysis.
for lumbar ADR; and 3 moderate quality randomized controlled trials, 22 low quality case series, and 1 economic analysis for cervical ADR. Using a formal, objective method of evaluating evidence, the evidence based technology assessment report concluded that there was: no evidence comparing ADR to non surgical alternatives; moderate evidence of equivalent or superior short term efficacy of ADR compared to fusion; no evidence for broad safety conclusions, and moderate evidence of comparable short term safety profiles between ADR and fusion, and no evidence on cost effectiveness.

On October 17\textsuperscript{th}, 2008, the HTCC, an independent group of eleven clinicians, met at an open public meeting to decide on whether state agencies should pay for artificial disc replacement in the lumbar and cervical spine based on whether the evidence report and other presented information proves it is safe, effective and has value. The HTCC reviewed the report, including peer and public review comments; and invited and heard public comments at the meeting. Meeting minutes detailing the discussion are available through the HTA program or online at \url{http://www.hta.hca.wa.gov} in the committee section.
B. Committee Findings

Having considered the evidence based technology assessment report and the written and oral comments, the committee identified the following key factors and health outcomes, and evidence related to those health outcomes and key factors:

B.1. Evidence availability and technology features

The committee finds the following key factors relevant to the coverage decision:

1.1. Five (two lumbar and three cervical) randomized controlled studies form primary evidence base comparing ADR to fusion surgery. Efficacy of comparator treatment, fusion remains uncertain and given evidence uses only this comparator, it limits the ability to fully answer efficacy/effectiveness question.

1.2. No evidence compares ADR to optimal medical treatment.

1.3. The RCTs were primarily conducted for FDA approval and were designed to prove that the new treatment is no worse than the comparator (non-inferiority design). Studies were not blinded, though this remains a difficulty of most surgical trials.

1.4. FDA trial “success” is defined based on specified clinical outcomes that must be within a margin to be not worse than the alternative. FDA specified success focus on clinical or surgical success (e.g. devise operation (technical performance, no device failure, no deterioration) and an ODI improvement of 25%).

B.2. Is the technology safe?

The committee separately discussed lumbar and cervical ADR safety outcomes. The committee found the following key evidence on safety from the evidence report:

2.1. No case related deaths were reported for either lumbar or cervical ADR.

2.2. No data on long term safety and complication rates was available for either lumbar or cervical ADR.

2.3. FDA database (MAUDE) listed about 500 safety events, but does not include denominator information.

2.4. L-ADR device related failure that required reoperation, revision, or removal as reported in trials was not statistically different (fusion 2.7 and 8.1% vs. ADR 5.4 and 3.7%)

2.5. L-ADR short term complications from trials and case series varied greatly from 1% to 60% - heterotopic ossification, hematoma, subsidence, and new or residual pain, secondary fusion especially had high ranges. No statistical differences in major adverse events/complications from trials between fusion and L-ADR.

2.6. C-ADR device related complications and failures (2.9%) occurred statistically significantly less than with fusion (8.9%).

2.7. C-ADR short-term complications from trials showed similar adverse events where the differences were not statistically significant (e.g. 26.4% vs 24.9% serious adverse events). Rates of complications from case series varied broadly (dysphagia 0% to 100%; new or residual pain 1% to 33%). No denominator information for Maude safety events.

B.3. Is the technology effective?

The committee found that there were multiple key health outcomes that were significant in assessing the technology’s effectiveness. The report identified the following evidence:
3.1. Neither lumbar nor cervical ADR had evidence on the long term durability of the device and efficacy of the intervention.

3.2. No studies looked at subgroups or performed subpopulation evaluation to determine those most or least likely to benefit. Applicability in older population and generalizability outside trial population unclear.

3.3. Return to Work and quality of life measures were not adequately reported on in either lumbar or cervical studies.

3.4. Preservation of flexibility measure was reviewed. This is generally not a comparative measure with fusion because fusion is designed to limit motion. Trials demonstrate pre operative motion generally maintained with ADR; both C-ADR and L-ADR had greater motion preservation than fusion.

3.5. A key proposed health benefit of ADR over fusion is that preservation of motion will relieve adjacent level stress/adjacent segment disease (ASD). This outcome was not measured in the randomized controlled trials. For L-ADR, non-randomized studies reported ASD in 0% to 34%. For C-ADR, ASD was reported at 1% in C-ADR vs 3% in fusion in RCT; other studies reported ASD rates of 1% to 7%.

3.6. Pain Relief. The evidence report concluded that ADR appears to provide as good or greater pain relief for single level disease than fusion. VAS pain score reductions for patients receiving L-ADR, over 2 years, were statistically significant. For C-ADR, both surgical groups reported clinically significant pain relief. There were no statistical differences in pain relief between C-ADR and fusion as measured out to two years.

3.7. Functional improvement. The evidence report included analysis on SF-36, clinical success and ODI for L-ADR. SF-36 a common health survey, scores demonstrated higher improvement on physical and mental component with L-ADR over fusion at 12 months (81% versus 77%). The clinical success (FDA measures) including ODI improvement, pooled at 57% improvement for fusion and 65% for L-ADR. For C-ADR, the primary measure used was neck disability index (NDI). NDI improvement in score of at least 15 points reached in both groups - 80% fusion and 82% C-ADR. The difference was not statistically significant.

3.8. Neurological success for C-ADR was defined in the trials to include both maintain and improve neurological function. 78% of C-ADR and 67% of fusion patients achieved bar – a statistically significant difference.

3.9. A composite measure of overall clinical (surgical) success defined by FDA standard – 66% C-ADR success vs. 55% fusion success.

3.10. Patient Satisfaction measured in two lumbar ADR trials and one cervical trial was reported higher for ADR than for fusion and more ADR patients than fusion patients would choose their treatment again. Not reported when the patients were asked questions; variable tools used.

B.4. Is the technology cost-effective?

The committee found that there was key information about cost and value:

4.1. The evidence report summarized two technology assessments that did include economic analysis comparing fusion and L-ADR (Ontario and Australia). These resulted in mixed findings that may suggest L-ADR has similar costs to fusion, but finding was not supported in Ontario analysis and could be dependent on fusion.
procedure used. One Australian HTA concluded that C-ADR and fusion costs were the same. Analysis includes assumptions related to health care system; practice patterns, and reimbursement mechanisms not present in US.

4.2. Economic studies reflected short time horizons to assess the potential cost-effectiveness of ADR technology and need appropriate comparator.

4.3. Approximate cost for ADR in WA based on 50% of hospital costs: L-ADR at $20,113 and C-ADR at $14,344.

4.4. No manufacturer provided any cost data.

C. Committee Conclusions

Having made findings as to the most significant and relevant evidence regarding health outcomes, key factors and identified evidence related to those factors, primarily based on the evidence based technology assessment report, the committee concludes:

C.1. Evidence availability and technology features

The committee concludes that the best available evidence on artificial disc replacement has been collected and summarized.

1.1. There is moderate evidence from 5 randomized controlled trials and about 40 uncontrolled studies about several important health outcomes for artificial disc replacement. The randomized trials have shared limitations: some methodological flaws, fusion as only comparator, non-inferiority design, lack of long term data, and measure/definition of success.

1.2. The controlled studies compare surgical options only. Fusion surgery as a treatment for spine pain is still not established a clearly superior option, so the lack of inclusion of optimized medical management severely limits the results.

1.3. As compared to fusion, a currently approved alternative, the overall evidence is moderate and demonstrates at least equivalence of ADR in short term safety and efficacy.

1.4. Longer follow up data, especially around safety events and reoperation rates is needed (often this evidence comes from non RCT data such as registries). Also, the post approval FDA studies requiring up to seven year follow up should be monitored.

C.2. Is it safe?

The committee concludes that the comprehensive evidence reviewed shows that the technology has been proven at least equally safe as a currently offered alternative, fusion. Key factors to the committee’s conclusion include:

2.1. Moderate evidence demonstrated that L-ADR has a similar safety profile as lumbar anterior or circumferential fusion two years following surgery. Longer term safety on L-ADR is not known.

2.2. Moderate evidence demonstrated that C-ADR tends to be safer than fusion as measured by the risk of device failure and surgical complications up to two years following surgery. Longer term safety on C-ADR is not known.
C.3. **Is it effective?**

The committee concludes that the comprehensive evidence reviewed shows that the technology has been proven equally or more effective as a currently offered alternative, fusion. Key factors to the committee’s conclusion include:

1. While there is no evidence comparing ADR with non-operative care, there are five moderate quality, controlled studies comparing ADR with a currently performed alternative, fusion. Based on the limited comparator and other evidence limitations, the evidence of efficacy should not be generalized beyond carefully selected patients that match trial and FDA indications.

2. Moderate evidence demonstrated that the efficacy/effectiveness of L-ADR is comparable with fusion up to two years following surgery based on a composite measure for FDA approval of overall clinical success, pain improvement, an ODI and SF-36 improvement.

3. Moderate evidence demonstrated that the efficacy/effectiveness of C-ADR is equal to fusion for pain and function and potentially superior to fusion for neurological and overall success up to two years following surgery.

4. There is insufficient evidence to draw conclusions regarding the safety and efficacy of ADR in special populations or populations outside those studied for FDA approval. Thus, coverage should be limited to studied indications.

C.4. **Is it cost-effective?**

The Committee concludes that the comprehensive evidence review does not show that the technology is more cost effective. Although cost-effectiveness was not a major decision factor, the committee concluded cost-effectiveness is unproven because of insufficient evidence.

1. The cost analyses were limited by short time horizons, comparators chosen, and differences with US health system, and provided mixed answers. For L-ADR, one assessment showed an increase in cost based on the device cost and another showed similar or possibly reduced cost based primarily on shorter hospital stays for L-ADR. For C-ADR, one cost analysis showed similar surgical costs, but higher total cost with C-ADR due to device cost.

C.5. **Medicare Decision and Expert Treatment Guidelines**

The committee deliberations included a discussion of National Medicare Decisions and expert treatment guidelines, and an understanding that the committee must find substantial evidence to support a decision that is contrary. RCW 70.14.110. The independent evidence report identified a national medicare coverage decision on lumbar fusion and no expert treatment guidelines. The committee’s conditional coverage is consistent with the national medicare decision to not cover lumbar ADR for patients older than 60 years of age.

**D. Committee Decision**

Based on the deliberations of key health outcomes, the committee decided that evidence on Artificial Disc Replacement demonstrates net health benefit because of moderate level evidence based on randomized controlled trials of effectiveness and short term safety.
The committee found that artificial disc replacement, as compared to fusion, was proven to be equally or more safe and effective, and the cost was not a significant factor for this decision based on inconclusive data. Based on these evidentiary findings, the committee voted for conditional coverage as follows:

- Lumbar-ADR: 6 cover with conditions and 2 no coverage
- Cervical-ADR: 8 cover with conditions

**E. Health Technology Clinical Committee Authority**

Washington State’s legislature believes it is important to use a scientific based, clinician centered approach for difficult and important health care benefit decisions. Pursuant to chapter 70.14 RCW, the legislature has directed the Washington State Health Care Authority, through its Health Technology Assessment program to engage in a process for evaluation process that gathers and assesses the quality of the latest medical evidence using a scientific research company and takes public input at all stages. Pursuant to RCW 70.14.110 a Health Technology Clinical Committee (HTCC) composed of eleven independent health care professionals reviews all the information and renders a decision at an open public meeting. The Washington State Health Technology Clinical Committee (HTCC), determines how selected health technologies are covered by several state agencies. RCW 70.14.080-140. These technologies may include medical or surgical devices and procedures, medical equipment, and diagnostic tests. HTCC bases their decisions on evidence of the technology’s safety, efficacy, and cost effectiveness. Participating state agencies are required to comply with the decisions of the HTCC. HTCC decisions may be re-reviewed at the determination of the HCA Administrator.

FDA Approved Devices: Use Indications and Contra-Indications


Conditions for Lumbar ADR (L3-S1 only)

A. Failed 6 months of conservative management (FDA)

AND

B. DDD at single level from L3-S1 (Prodisc-L) or L4-S1 (Charite) (FDA)

AND

C. NO spondylolisthesis > Grade 1, no radiculopathy or disc herniation, no spinal stenosis, no pars defect (FDA)

AND

D. Failure or inability to access a structured, intensive, multi-disciplinary program. HTA decision)

Conditions for Cervical ADR (C3-C7 only)

A. Single level intractable radiculopathy (symptomatic nerve root compression) or myelopathy (symptomatic spinal cord compression) as determined from history, physical exam, and imaging studies (FDA)

AND

B. For reconstruction of the disc following single level discectomy for condition A

AND

No marked instability on flexion/extension x-rays (>3 mm translation or >11 deg angular rotation) OR No severe spondylosis characterized by bridging osteophytes or a loss of disc height >50% or an absence of motion (<20) (FDA)
January 6, 2009

Ms. Leah Hole-Curry  
Program Director  
Health Technology Assessment Program  
676 Woodland Square Loop SE  
Olympia, Washington 98504-2712

RE: HTA Artificial Disc Replacement Draft Decision Comments

Dear Ms. Hole-Curry:

We appreciate the opportunity to comment on the Draft Health Technology Committee Findings and Coverage Decision for Artificial Disc Replacement. As you are probably aware, Medtronic Spinal and Biologics Division manufactures products that treat a variety of disorders of the spine. These products are utilized by spinal and orthopedic surgeons to treat patients and restore their quality of life. As the manufacturer of cervical discs, we are very interested in the draft decision on cervical disc arthroplasty.

We applaud the draft decision allowing cervical disc arthroplasty in accordance with FDA approved indications. However, we do have concerns regarding the limitation of coverage by requiring completion of a “structured, intensive, multi-disciplinary program for management of pain, if covered by the agency.” This program, entitled “SIMP” by the state agencies, is being created as a condition of coverage for the lumbar fusion review that the HTCC conducted in November 2007. A review of the recordings made at the hearing for artificial disc replacement, does not provide evidence that this limitation was established for cervical discs. The HTCC motion stated that FDA indications must be met and that a patient must not have the contraindications established by the FDA. The group also stated that staff should be directed to conduct a feasibility study on the creation of a Registry to track disc arthroplasty usage like they are creating for lumbar fusion.

There was no mention in the hearing minutes of creating a SIMP program for cervical disc arthroplasty and we recommend that the SIMP program not be utilized as a condition for coverage of cervical disc arthroplasty for numerous reasons.

First, the SIMP program being created by the agency medical directors, with input from the IMAC, is for lumbar fusion and would not necessarily be appropriate for cervical disc arthroplasty. Therefore, a separate SIMP program would have to be created for cervical disc arthroplasty.

When Life Depends on Medical Technology
Second, the SIMP program for lumbar fusion is still being created 14 months after the lumbar fusion decision. This is an inappropriate time period for creation of coverage criteria associated with an HTCC decision. Additionally, patients should be able to take advantage of the newest technology after a decision has been made without waiting an undue amount of time, due to development of coverage criteria (e.g. the SIMP program).

Finally, and due to the first two issues above, the conservative care program set as a condition for coverage of cervical disc arthroplasty should be established by the surgeon caring for the patient. Surgeons who have been trained in disc arthroplasty are the best resource to know when their patients have exhausted appropriate conservative care and are candidates for a cervical disc. Medtronic recommends that the HTCC defer to the expertise of the spine surgeon and allow them to make the determination as to when conservative care has failed and surgical intervention is required.

After the thorough review conducted by the HTCC, coverage decisions should not be unduly delayed by additional conditions. We would encourage the HTCC to recognize the practical application that the decisions have on patients and ensure that appropriate care is not delayed by the creation of a conservative care program. This issue is made even more important because of the excessive time it has taken to create a conservative care program for the HTCC’s lumbar fusion decision 14 months ago.

Thank you again for the opportunity to comment on the Artificial Disc Replacement Findings and Coverage Decision Draft. We are more than happy to respond to any questions you have regarding these comments.

Sincerely,

Dena Scearce
Director, State Government Relations
Medtronic Spinal and Biologics

cc: Ms. Denise Santoyo
Dear Washington State Health Technology Assessment Program,

The Multisociety Spine Work Group would like to thank the HTA for the opportunity to participate in the October Clinical Meeting on the topic of artificial discs and commends the Clinical Committee for their review and resulting decision. The Work Group believes this decision will ultimately be of benefit to both Washington State and patients alike.

Sincerely,

James R. Bean, MD
American Association of Neurological Surgeons

Thomas A. Zdeblick, MD
Cervical Spine Research Society

P. David Adelson, MD
Congress of Neurological Surgeons

Charles Branch, MD
North American Spine Society

Oheneba Boachie-Adjei, MD
Scoliosis Research Society

Karin Büttner-Janz, MD, PhD
Spine Arthroplasty Society

Pamela M. Hayden
Director, Research & Quality Improvement
North American Spine Society
8320 St. Moritz Drive
Spring Grove, IL 60081
815.675.0021
Fax 815.675.3137
January 6, 2009

Washington State Health Care Authority  
c/o Leah Hole-Curry, Program Director,  
Health Technology Assessment Program  
Health Technology Assessment Committee  
P.O. Box 42712  
676 Woodland Square Loop SE  
Olympia, WA 98504-2712

Re: Re-Review of Upright MRI Technology

Dear Health Technology Clinical Committee:

Please accept this letter as a formal request that the Health Technology Clinical Committee (HTCC) place the upright MRI technology for re-review on its 2009 agenda. This request is made pursuant to my conversation with Leah Hole-Curry, Program Director, Health Technology Assessment Program. The upright MRI technology was originally reviewed by the HTCC in 2007. In May 2007 the HTCC made a determination that the upright MRI would not be covered.

Since the May 2007 determination, additional studies and reports have been conducted and issued. I have attached for your review a copy of our October 22, 2008, letter sent to the Washington State Health Care Authority (HCA) requesting that the Administrator place the upright/positional MRI on the 2009 agenda. On December 12, 2008, the HCA Administrator issued a notice of selected health technologies to be reviewed in 2009, by the HTCC. The upright MRI was eligible for re-review but not placed on the 2009 agenda.

We disagree with the Administrator’s decision to not place the upright MRI on the 2009 agenda. We ask that the HTCC independently place the upright MRI on the 2009 agenda for re-review.

We ask that time be set on your February 6, 2009, meeting agenda to address this request. Please let me know what further information you may require to make your assessment. In addition, we will make available, at the February 6, 2009, meeting, individuals to answer any questions that you may have regarding the technology and/or the new studies.
Thank you for your consideration of this request. Please let me know if you have any questions or require further information.

Very truly yours,

LANE POWELL PC

Robert A. Battles

RAB/tlm
Enclosure
cc: Mr. Peter Solodko
122037.0002/34685.1
October 22, 2008

Washington State Health Care Authority  
Health Technology Assessment Committee  
P.O. Box 42712  
676 Woodland Square Loop SE  
Olympia, WA 98504-2712

Re: Re-Review of Upright MRI Technology

Dear HTA Committee:

Our office represents Capital Imaging, LLC.

We are aware of the Preliminary Recommendations for 2009 HTA Technology Topics dated October 15, 2008, posted on the Health Care Authority/HTA website that indicates under the Re-Review Recommendations section that “[a] PubMed literature scan was completed by the HTA clinical consultant . . . [that] did not reveal any significant new evidence that might lead to a different conclusion.” However, no mention of what information may have been reviewed or obtained and considered by the HTA.

We disagree the HTA’s recommendations and submit that additional significant information is and has been available during the re-review period. Attached is the following information that was not previously available during the review process and prior to the HTA Committee’s determination in May 2007.


2. Letter from Jeffrey C. Wang, M.D., Chief, Orthopaedic Spine Service, UCLA Comprehensive Spine Center and Associate professor of Orthopaedic and Neurosurgery, UCLA School of Medicine, regarding indications for positional / kinetic MRI, dated August 6, 2007.

Capital Imaging therefore requests that the Upright MRI technology be placed on the 2009 HTA Technology Topics agenda. In the alternative, Capital Imaging requests the HTA perform a re-review of the Committee's determination of May 18, 2007, and consider the enclosed materials as part of its re-review determination process.

Very truly yours,

LANE POWELL PC

Robert A. Battles

RAB:ms
Enclosures
cc: Mr. Peter Solodko
122037.0002/34184.1
ATTACHMENT 1
SIC 1: Diagnostic Imaging: The Pros and Cons of Static, Dynamic, Upright and Inline Spine Imaging. [Official Program Schedule, Wednesday, October 24, 2007]

Room 128

Moderator: Christopher B. Chaput, MD

Spinal surgery is changing from a discipline that uses primarily static imaging of the neuraxis to define pathology to one where dynamic studies are increasingly available. The need for evidence-based information on these imaging modalities is expected to increase as more options for noninvasive instrumentation become available to the spine surgeon. Discussion will center around currently available dynamic and static imaging and will seek to explore the relationship of such imaging to diagnosing and treating patients.

B: The Effect of Lumbar Flexion and Extension on the Central Canal With Dynamic MRI

Feng Wei, MD\(^1\), Soon Woo Hong, MD\(^1\), Jun Zou, MD\(^1\), Benjamin Yus, MD\(^2\), Masahito Miyazaki, MD\(^3\), Yuhichiro Marichida, MD\(^3\), Abhijit Alamgir, MD\(^3\), Jean-Joseph Abbed, MD\(^4\), Richard Weng, MD\(^4\), LAA, CA, USA; \(^1\)CA, USA; \(^2\)University of California Los Angeles, Los Angeles, CA, USA; \(^3\)University of California, Los Angeles, CA, USA; \(^4\)University of California, San Diego, CA, USA

BACKGROUND CONTEXT: Lumbar central canal stenosis is defined as the reduction in the diameter of the spinal central canal, which causes neurogenic claudication and radicular leg pain. Previous myelography and in vitro study of cadaveric specimens showed that extension of the spine caused protrusion of the intervertebral disk, bulging of the ligamentum flavum, and spurious osteophytes, resulting in a narrowing of the canal. However, few normative studies exist to show these results. Dynamic MRI studies can show with high precision the amount of change of the diameter of the spinal canal with flexion and extension of the spine.

PURPOSE: The purpose of this study was to define the diameter changes of the spinal canal at each level of the lumbar spine with dynamic MRI studies, to document the amount of change, and to see how progressive degeneration of the disc at the functional spinal unit will affect these values.

STUDY DESIGN/SETTING: This was a retrospective study on patients who presented with low back pain and were examined by dynamic MRI to determine the effect of lumbar flexion and extension on spinal canal.

PATIENT SAMPLE: Lumbar MR images for 119 patients, including 192 male and 169 female, (18-85 years of age), with low back pain, were obtained.

OUTCOME MEASURES: All radiological data on MRI was recorded on computer based measurement from MRI taken by flexion, neutral, and extension.

METHODS: All patients were examined in sitting flexion 40 degree, upright, and extension 10 degree with a 0.5 T dynamic MRI scanner. Quantitative measurements of canal diameter in the sagittal midline in disc level were obtained for each position. Degeneration was graded according to the grade of the discs in T2 weighted images. Change ratios of the canal diameter from neutral position to flexion or extension were calculated to reflect the extent of change relative to the grade of degeneration.

RESULTS: Statistically significant differences in canal diameter were obtained between neutral and flexion position and between neutral and extension position for L2-3 to L5-S1 levels. Results showed that flexion increased the canal diameter and extension decreased the canal diameter. Change ratio of L4-5 was greatest in both flexion and extension. In flexion, the change ratio positively correlated with the degree of degeneration in L2-3 to L5-S1. In extension, the change ratio negatively correlated with the degree of degeneration in L3-4 only.

CONCLUSIONS: Dynamic MRI can demonstrate spinal canal diameter changes in lumbar flexion and extension and also show the amount of change in the cross-sectional area with the highest accuracy. The spinal canal is widest in flexion and narrowest in extension. The relief of spinal stenosis in flexion is greater when the degree of degeneration is more severe. Furthermore, the less amount of degeneration, the greater the change in extension of the canal diameter. This study is the first to fully define the amount of diameter change of the spinal canal with flexion and extension of the spine, quantify the change at each level, and demonstrate how these values change with the increasing amount of degenerative grade of the disc at the functional motion segment.

FDA DEVICE/DRUG STATUS: Dynamic MRI: Approved for this indication.

doi: 10.1016/j.spinee.2007.07.096

NII: Positional MRI: A Valuable Tool in the Assessment of Cervical Disc Bulge

Payam Hassanzai, MD\(^1\), Soon Woo Hong, MD\(^1\), Masahito Miyazaki, MD\(^2\), Mark Ashkan, BS\(^3\), Jeffrey Weng, MD\(^4\), University of California, Los Angeles, Santa Monica, CA, USA

BACKGROUND CONTEXT: Positional MRI (pMRI) has recently been proposed as an alternative to conventional MRI techniques, pMRI offers the advantage of assessing cervical spine pathology in the neutral, flexion, and extension positions. pMRI also allows examination of the cervical spine in more physiologic, weight-bearing position as compared to traditional spine MRI imaging. A recent review of the literature demonstrated no studies to-date that have investigated the amount of cervical disc bulge in the neutral, flexion, and extension positions.

PURPOSE: The purpose of this study was to determine if adding flexion and extension MRI data to traditional neutral views would be beneficial in the evaluation of cervical disc bulges.

STUDY DESIGN/SETTING: Patients with cervical disc bulge signs and symptoms underwent pMRI in neutral, flexion, and extension. The images were analyzed using novel computer measurement technology to objectively quantify the amount of disc bulge.

PATIENT SAMPLE: One hundred eighty-three patients with cervical spine symptoms were included in the study. This represented 978 cervical discs in total. There were 69 males and 94 females. The mean age was 44.1 years (range 13-93).

OUTCOME MEASURES: Disc bulge was measured at the amount of extension of the disc beyond the intervertebral space. Discs with less than 2.0 mm disc bulge in the neutral position were selected and compared with their respective flexion and extension data.

METHODS: Disc bulge was measured using MRI Analyzer™ Version 3 (Tenetor Corporation, Bellflower, CA) anatomic software to objectively quantify the amount of disc bulge in millimeters. The statistical significance was calculated using the chi-square test.

RESULTS: The mean disc bulge was 1.96 mm in neutral, 1.86 mm in flexion and 1.93 mm in extension (p=0.078) discs. For discs with less than 2.0 mm disc bulge in neutral (n=539 discs), the results were as follows: 18.18% 2.0 mm bulge in flexion and 23.75% 2.0 mm bulge in extension (p=0.025). In addition, 2.41% 3.0 mm bulge in flexion and 3.34% 3.0 mm bulge in extension (p=0.30). Using 2.0 mm of disc bulge as a cutoff value, the false negative ratio for the neutral position alone compared to flexion and extension was 25.08%.

CONCLUSIONS: A significant increase in the degree of cervical disc bulge was noted by comparing flexion and extension views as compared to neutral views alone. This study also suggests that extension MRI views yield a higher detection rate of missed cervical disc bulges than flexion views. Flexion and extension MRI views provide a valuable, added information when assessing patients for cervical disc bulge. This data suggests that positional MRI might be especially beneficial in patients with symptomatic radiculopathy and unimpressive static MRI studies.

FDA DEVICE/DRUG STATUS: This abstract does not discuss or include any applicable device or drug.

doi: 10.1016/j.spinee.2007.07.097
BACKGROUND CONTEXT: Many people suffer from cervical degenerative disease. The degree of degeneration of cervical spine has not been compared with the extent of cervical spine mobility. The effect of degeneration of the disc on the overall motion of the functional spinal unit is also not definite. Dynamic Motion MRI studies can demonstrate the mobility of each motion segment and define where the motion of the spine occurs, and it can relate it to the grade of degeneration.

PURPOSE: To define the relationship between the grade of disc degeneration and the motion unit in cervical spine and to elucidate how the role of each cervical spine unit for flexion-extension motion changes with degeneration with dynamic MRI.

STUDY DESIGN/SETTING: Prospective patients with neck pain were enrolled and obtained a dynamic flexion-extension MRI of the cervical spine.

PATIENT SAMPLE: 168 patients were permitted to enroll our study with symptomatic neck pain with/without radiography or myelography. Methods: All radiologists on MRI were recorded on computer based measurement from MRI taken by flexion, neutral and extension. MRI Analyzer in true MRI which included 76 point marked in each image was undergone automatically all measurement and calculations with regard to translational motion and angular motion on each segment. According to grading system on the basis of the literature, two observers analyzing MRIs graded 5 (grade 1 to V) in each of intervertebral disc on the T2-weighted sagittal images.

RESULTS: On the each cervical unit, compare to more normal discs with Grade I and II mild degeneration, translational motion and angular motion were significantly increased for segments with discs with higher degenerative grades (Grade III and IV). However, the authors observed that the translational motion and angular motion of the segments decreased significantly in severe Grade V degeneration. For the Grade I and II segments, C4/5 and C5/6 units contributed the majority of the total angular mobility of the spine. For Grade III and IV degeneration, the segments of C3/4 and C6/7 units increased as well as C4/5 and C5/6 units. In Grade V, the roles of C4/5 and C5/6 units for total angular mobility decreased.

CONCLUSIONS: Following degeneration, the changes of translational motion and angular motion were observed. Namely, the authors demonstrated the changes that occur with progressive degeneration. The angular motion and translation moves from normal disc (Grade I and II) to a more unstable phase (Grade III and IV) to a more stable stage with more stability (Grade V). We also demonstrated the contribution of different levels to overall motion that occurs with degeneration.
ATTACHMENT 2
August 6, 2007

Cervical Spine
Indications for Cervical positional / kinetic MRI (kmMRI) as study of choice:
Neuroimaging recommendations are based on our study of 1,000 patients having kinetic MRI scans demonstrating over 20% false positive rate of instability, cervical stenosis, cervical disc herniations, cervical spondylolisthesis, and other missed pathologies:

1. Generalized cervical or neurological complaints (e.g., back pain or radiculopathy)
2. Patients with spondylolisthesis to characterize the amount of instability present.
3. Patients with C1-C2 instability to characterize the amount of spinal cord compression.
4. Patients with suspected spinal pathology after cervical trauma with or without evidence of abnormalities on standard studies to characterize the amount of instability and the injured structures.
5. Patients who are surgical candidates for cervical disc arthroplasty for pre-operative evaluation to confirm the absence of instability, facet pathology, or neuroforaminal narrowing.
6. Patients who are candidates for cervical fusion surgery with evidence of degenerative pathology to rule out adjacent level disease, instability, or stenosis.
7. Patients who are surgical candidates for posterior motion sparing devices (e.g., pedicle-based motion-altering rods)

Relative indications:

1. Pre-operative evaluation for patients undergoing cervical decompressive surgery to help decide on the appropriate levels for decompression.
2. Pre-operative evaluation for patients undergoing cervical fusion surgery to help decide on the appropriate levels for fusion.
3. Pre-operative evaluation for patients undergoing cervical surgery to rule out adjacent segment problems, characterize the amount of instability present, and the expectations from the surgical procedure through the range of motion of the cervical spine.
4. Patients with rheumatoid arthritis to rule out atlanto-axial instability.

Thoracic Spine
Indications for Thoracic positional / kinetic MRI (kmMRI):

1. Generalized thoracic or neurological complaints (e.g., back pain or radiculopathy).
Lumbar Spine
Indications for lumbar positional/motion MRI studies (kMRI):

Note: These recommendations are based on our study of 1,000 patients having kinetic MRI studies demonstrating over 20% miss rates of instability, lumbar stenosis, lumbar disc herniations, lumbar spondylolisthesis, and other missed pathologies.

Indications for Lumbar kMRI studies:

1. Generalized lumbar or neurological complaints (e.g., back pain or radiculopathy)
2. Patients with spondylolisthesis to characterize the amount of instability present.
3. Patients who are surgical candidates for lumbar disk arthroplasty for pre-operative evaluation to confirm the absence of instability, facet pathology, or neuroforaminal narrowing.
4. Patients who are candidates for lumbar fusion surgery with evidence of degenerative pathology to rule out adjacent level disease, instability, or stenosis.
5. Patients with spinal stenosis who are surgical candidates for interspinous spacer (to confirm/predict the amount of decompression from device and to confirm that they are appropriate candidates)
6. Patients who are surgical candidates for posterior motion sparing devices (e.g., pedicle-based motion-allowing rod)

Relative indications:

1. Pre-operative evaluation for patients undergoing lumbar decompressive surgery to help decide on the appropriate levels for decompression.
2. Pre-operative evaluation for patients undergoing lumbar fusion surgery to help decide on the appropriate levels for fusion.
3. Pre-operative evaluation for patients undergoing lumbar surgery to rule out adjacent segment problems, characterize the amount of instability present, and the expectations from the surgical procedure through the range of motion of the lumbar spine.

Follow-Up Study
Note: Performing MRI of the spine (C/F) as a follow-up study is recommended in cases where there is evidence of symptomatic or related compressive pathology on standard recent MRI studies (e.g., if the treating physician believes that the standard recent MRI missed clinically important pathology).

If you have any further questions, please do not hesitate to contact me.

Sincerely,

[Signature]

Jeffrey C. Wang, M.D.
Chief, Orthopaedic Spine Service
UCLA Comprehensive Spine Center
Associate Professor of Orthopaedic and Neurosurgery
UCLA School of Medicine
1250 16th Street, 7th Tower, #745
Santa Monica, CA 90404
Tel: (310) 319-3334 or 319-3827, Fax: (310) 319-5055
ATTACHMENT 3
of HBJQ includes: S62-72 and S8-12. Spearman's Rank Order Correlation was used to determine the relationship between individual radiographic parameters and HBJQ. Pedestrians were then grouped into dichotomized deformity patterns (including: High [H1-12] Low grade [0.11] by segmenting Cervical, kyphosis or lordosis at L5-S1, and high [>30 degrees] and low [<30 degrees] pelvic tilt). Dichotomous variables were then analyzed using Student's t-test.

RESULTS: 27 adults (age 10-61) with spondylolisthesis had complete radiographs and clinical data. There was a moderate correlation between C7 cervical balance and appearance (r=0.7), and slighty (r=0.6) on the S2-5 lumbar spine. All other radiographic parameters had a low independent correlation with HBJQ. Grouping patients by low or high kyphosis or lordosis at L5-S1, adults with a normal or kyphotic angle at L5-S1 had significantly more pain (p=0.01), functional limitations (p=0.02), and manual health complaints (p=0.01) than patients with a kyphotic angle. Adults with a high pelvic tilt had significantly worse scores for appearance (p=0.05), pain (p=0.05), function (p=0.02), and physical components summary (p=0.02). Adults with a high grade of disc herniation (MIV) had more variation in physical role (p=0.04) and mental health (p=0.02) than those with a low grade of disc herniation.

CONCLUSIONS: Surgical strategies for the management of high-grade spondylolisthesis require a detailed understanding of patient-specific factors. Factors that influence high-grade spondylolisthesis may cause lumbar and pelvic misalignment with the resulting symptoms of high-grade spondylolisthesis.

OUTCOME MEASURES: Cervical incisional widening was defined by the distance between sphenoid process tips in full flexion versus flexion distance in full extension. The extent of the superior extension of the C4 vertebrae.

METHODS: Each cervical flexion-extension x-ray was captured with an x-ray fluoroscopy imaging system (CMAX, Medical Imaging, Inc., Boston, MA). The radiology and stereolithography software (CMa, Medical Imaging) allows the specific vertebral to be superimposed in flexion and extension. This software has been validated to measure intervertebral motion with errors less than 0.8 degrees and interobserver translational motion with errors less than 0.5 mm.

RESULTS: Of the 156 asymptomatic subject entered into the study, 7 had lesions that could not be traced, and the final level could be analyzed due to problems visualizing the anatomy. The following subjects provided 65 cervical levels for analysis. The mean, median, and the lower and upper limits of the 95% confidence interval for each cervical level are provided in Table 1. The lower cervical spine contributed the most to motion. A comparison of the motion at each given level to its adjacent levels showed significantly more motion than its neighbor (Figure 1).

CONCLUSIONS: This data supports the general rule that intervertebral motion in the lower cervical spine is less than 20% of that at the adjacent levels representing an abnormal finding and should raise clinical concern for disease or injury. This data may be used in the planning of surgery and in the assessment of outcomes in the adult.

FOA DISCLOSURES/STATUS: This abstract does not disclose or indicate any clinical concerns or drugs.

Table 1. Summary of Data

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Figure 1. Means minimum absolute difference and 95% CI between adjacent levels in sphenoid process tip displacement.

doi: 10.1016/j.spinee.2007.01.005

Thursday, October 25, 2007
4:15-5:03 PM
Concurrent Session 2: Diagnostics

Azeem Mehmood, MD, John Young, MD, Palak Banerji, MD, Charles Callahan, MD, Houston, TX, USA

BACKGROUND CONTEXT: Lateral flexion and extension radiographs of the cervical spine are commonly used to evaluate injuries or disease that may not be apparent on static lateral files. Computer assisted imaging techniques are now commonly used in the analysis of cervical motion, and reference data for interpreting intervertebral motion and measurements have been published. However, there are currently no clinically validated criteria for assessing intersegmental process widening in flexion-extension x-rays of the cervical spine.

PURPOSE: The purpose of this study is to document normal intervertebral process widening in a cohort of asymptomatic adults, using previously validated computer-assisted tracking software.

STUDY DESIGN/SETTING: Prospective cervical flexion and extension radiographs were taken, the images traced, and intersegmental widening calculated for each cervical level.

PATIENT SAMPLE: 156 asymptomatic subjects were recruited to participate in the study. Patients were not recruited if they reported prior neck pain which was consistent with a visit to a physician.

REFERENCE/STATUS: This abstract does not disclose or indicate any clinical concerns or drugs.

Table 1. Summary of Data

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Figure 1. Means minimum absolute difference and 95% CI between adjacent levels in sphenoid process tip displacement.

doi: 10.1016/j.spinee.2007.01.005
Journal of the Neurological Sciences

Epidurals and Disc Herniation

BY

Robert J. Wiltse, M.D.

ABSTRACT

This paper reviews the current status of epidural injections in the management of low back pain. The indications, techniques, and results of epidural injections are discussed. The role of epidural injections in the evaluation of low back pain is also considered.

INTRODUCTION

Epidural injections have been used for many years in the treatment of low back pain. The technique has evolved over the years, and current indications for epidural injections are well established. The purpose of this paper is to review the current status of epidural injections in the management of low back pain.

INDICATIONS

The primary indication for epidural injections is the treatment of low back pain. Other indications include the evaluation of low back pain, the treatment of sciatica, and the treatment of radicular pain.

TECHNIQUE

The technique of epidural injection involves the injection of a local anesthetic and/or corticosteroid into the epidural space. The level of injection is determined by the location of the pathology.

RESULTS

The results of epidural injections vary depending on the indications and the technique used. In general, epidural injections are effective in the treatment of low back pain and sciatica.

CONCLUSIONS

Epidural injections are an effective treatment for low back pain and sciatica. They are also useful in the evaluation of low back pain. Further research is needed to determine the optimal indications and techniques for epidural injections.

FURTHER READING


null
PURPOSE: The purpose of this study was to compare the safety and accuracy of pelvic screw placement in the thoracic spine using the free-hand anterior technique with and without the knowledge of the pre-operative CT scan parameters.

STUDY DESIGN/SETTING: This study is a prospective, randomized, radiographic review of 22 patients.

PATIENT SAMPLE: 22 patients undergoing spinal deformity surgery were randomized into two groups. The average age was 14.6 years (10-18 years). All of the patients had idiopathic scoliosis, except one who had Scheuermann's kyphosis.

OUTCOME MEASURES: Pre-operative CT scans were obtained and used to assess pelvic screw placement. Posterior correction was performed and assessed as projecting medially into the intervertebral space, without evidence of pedicle violation or fracture. A total of 21 patients were included in the scoliosis and kyphosis groups.

METHODS: The average age was 14.6 years (10-18 years). All of the patients had idiopathic scoliosis, except one who had Scheuermann's kyphosis. The average Cobb angle was 103° (47-77 degrees). A total of 169 pelvic screws were placed. In group A, 18 screws (12.5%) were misplaced out of a total of 158 placed. In group B, 31 (19.4%) were misaligned out of 212 placed. Evaluation of pre-operative CT scans did not significantly affect the percentage of screws that were needing correction.

CONCLUSIONS: As practiced hands, there is not a significant difference in the incidence of pelvic screw misalignment based on the ability to perform pre-operative evaluation of CT scan-based pedicle parameters. With rigorous free-hand anterior technique, pelvic placement of screws is possible and can be done safely without the reliance on routine pre-operative CT. This can decrease a patient's total radiographic exposure over the course of treatment of their spinal deformity.

FDA DEVICE/DRUG STATUS: This abstract does not discuss or include any applicable devices or drugs.

doi: 10.1016/j.spinee.2007.07.169

BACKGROUND CONTEXT: Spinal deformities are diagnosed as an outcome of spinal deformity. This study evaluated the effectiveness of using X-rays in spinal deformity.

STUDY DESIGN/SETTING: One spinal deformity study was performed on various X-ray methods to determine the accuracy of using X-rays to correct spinal deformity.

PATIENT SAMPLE: 510 patients who were enrolled in this study and underwent spinal deformity correction.

OUTCOME MEASURES: All spinal deformity data was recorded as computerized measurements from MRI taken by Scoliosis, cervical, and lumbar MRI.

METHODS: MRI patients (166 female, 304 male, mean age 42 years, range 16-85 years) had taken dynamic MRI for the study. Twenty-three patients were selected to determine if the MRI was useful in spine deformity correction. The MRI was simply read on a normal or abnormal for facet arthrosis.
Friday, October 26, 2007
4:16-5:16 PM
Special Interest Poster Presentation 3:
Lumber - Non-fusion

146. Indirect Decompression (X-Stop) versus Conventional Decompressive Surgery for Lumbar Spinal Stenosis: A Prospective Randomized Trial
Bilal Bashir, MD, MPH, Steven Burg, MD, Paul Goetzinger, MD, PhD, Roger Johnson, MD, MPH, Erik Kristiansson, MD, PhD, Zbigniew Sadowski, MD, PhD, Landstingets Sjukhus, Sweden; Landstingets Sjukhus, Malmo, Sweden; Stockholm Spina Center, Upplands Väsby, Sweden

BACKGROUND CONTEXT: Although generally successful, decompressive surgery for lumbar spinal stenosis has its complications and requires hospitalization and rehabilitation. Furthermore, at failure rates for laminectomy and fusion for lumbar disc herniation in current series. Our study aimed to compare the clinical outcome of patients treated with the X-Stop device with those treated with conventional decompression.

PURPOSE: To compare the outcomes in terms of function, quality of life and re-operations after lumbar decompression versus conventional decompression for LSS.

STUDY DESIGN/SETTING: Prospective randomized study including patients with central spinal stenosis according to MRI or CT, eligibility to conservative treatment and willingness to participate in the RCT. Outcome at 6, 12 and 24 months.

PATIENT SAMPLE: 50 patients in each group, were randomly assigned to either X-Stop treatment or surgical decompression. No surgical treatment at the spine center in Sweden.

OUTCOME MEASURES: Function and satisfaction according to the Zurich Spinal Symptoms Questionnaire, quality of life according to SF-36 and incidence of repeat operations as well as complications. Pain on VAS scale.

METHODS: After completing all necessary follow-up, patients were randomized to either X-Stop treatment under local anesthesia or decompressive surgery in general anesthesia on 1 or 2 levels.

RESULTS: Follow-up data were available for complete 2 years of spinal sources for all patients in the X-Stop group and 6 months in the decompression group. Follow-up in the decompression group was performed at 6 months, but 24 months had similar outcomes as regards the SF-36 and the ZSQ and both groups were significantly improved compared with baseline.

CONCLUSIONS: Preliminary results demonstrate that, when compared, X-Stop decompression surgery may yield similar results as decompressive surgery in terms of function and quality of life. An increased risk of secondary surgery is obvious and will be analyzed regarding when the follow-up is complete.

FDA DEVICE/DRUG STATUS: X-Stop: Approved for this indication.

DOI: 10.1016/j.spinee.2007.07.171

147. Interscalene Dye Transplantation: A New Dimention in the Treatment of Degenerative Sling Diseases
Di-hsuan, MD, Shih Hsueh, MD, Yi Hsu, MD, Yi-Ju, Keith L. Liao, MD, Department of Orthopedic Surgery & Traumatology, The University of Hong Kong, Hong Kong, China

BACKGROUND CONTEXT: Sling fusion is a well-recognized technique for the treatment of disc disease. However, over compression is accelerated when lumbar decompression and fusion at the disc level have been performed. The authors have previously reported success using interscalene dye, autograft, fascia lata, and fresh frozen allografts as a non-fusion strategy in a clinical study.

PURPOSE: To review the process of development of the concept of interscalene disc allograft transplantation from a basic experiment in a pig to human trials, to demonstrate the feasibility and long-term clinical results of this innovative surgical option to the literenity.

STUDY DESIGN/SETTING: A prospective, non-blinded study of 20 patients who received a fresh-frozen allograft interscalene disc transplantation in the cervical spine.

PATIENT SAMPLE: Five patients, 4 males and 1 female, average age 59 years.

OUTCOME MEASURES: Pain scores and VAS scores were used to evaluate pain reduction. Serial MRI, plain and dynamic radiographs were used to monitor the status of the allograft.

METHODS: All patients with cervical disc herniation underwent anterior intervertebral disc excision, and the interscalene disc allograft transplantation. The mean follow-up period was 3 years and 3 months.

RESULTS: The average operating time was 1.5 hours and blood loss was 63 ml. At a minimum follow-up of 3 years, all patients had improvement of the preoperative myelopathic or radiculopathic symptoms. Good union between the recipient bone and the graft endplates was seen after two months with no graft migration or subsidence. No neurological improvement of preoperative symptoms was found. There was reduction of the disc height in the early postoperative period and mild degenerative changes at the final follow-up.

CONCLUSIONS: The mobility and stability of the allograft segment were preserved. Fresh-frozen allograft interscalene disc transplantation has been successfully performed in a clinical setting without neurological
OFFICIAL PROGRAM SCHEDULE

5:25–6:25 p.m.
Nursing Track SIG: Tissue Banking
Room 12B
Moderator: Beverly Pericuto, RN, BSN
This session focuses on bone graft physiology, allograft procurement and The Joint Commission's requirements.

Upon completion of this session, participants should be able to:
- Review bone graft physiology.
- Define allograft procurement methods.
- Identify The Joint Commission's requirements for handling of tissue.
- Discuss the hospital's required documentation for allograft use.

PA/MP Track SIG: Discussion of the Day's Educational Offerings
Room 16A
Moderators:
Ian W. Marks, MSc, PA-C
James Brezina, MD

Followed by:
6:30–8:00 p.m.
PA/MP Surgeon Reception
Ballroom D Foyer

Rehabilitation Professional SIGs: Where Are We as Perioperative Providers vs. Nonoperative Care Providers?
Room 18B
Moderators:
Michael L. Reed, DPT, OCS
Christopher J. Standaert, MD
Kenneth S. Yonehara, MD

Symposium SIG: The New Performance Measurement Program: Your Care will be Assessed
(See symposium description on page 38)
Room 15
Moderator: David A. Wong, MD, MSc, FRCSC(C)

Symposium SIG: Treatment of the Symptomatic Lumbar Degenerative Disc Disease: Repair, Regenerate and Rehabilitate (See symposium description on page 41)
Room 17
Moderator: J. Kenneth Burkus, MD

SIG 1: Diagnostic Imaging: The Pros and Cons of Static, Dynamic, Upright and Supine Imaging
Room 12B
Moderator: Christopher D. Chaput, MD

Spine surgery is changing from a discipline that uses primarily static imaging of the neuraxis to define pathology to one where dynamic studies are increasingly available. The need for evidence-based information on these imaging modalities is expected to increase as more options for nonfusion instrumentation become available to the spine surgeon. Discussion will center around currently available dynamic and static imaging and will seek to explore the relationship of such imaging to diagnosing and treating patients.

SIG 2: Why Worry about Conflict of Interest?
Room 16B
Moderator: Wilton Bunch, MD

The term "conflict-of-interest" is being used in multiple medical settings and is discovered or implied somewhere nearly every day. It is easy to become overly saturated and ignore the issue. But, the issue is very real. This SIG discusses why it is important for every physician to be aware of and understand real and potential conflicts-of-interest.

5:25–6:25 p.m.
SIPP 1: Biologics
Room 12A
Moderator: Louis Jenis, MD

5:25–5:50 p.m.
26. Increased Expressions of Nerve Growth Factor and Its Tropomyosin-related Kinase A Receptor: A Potential Mechanism for Development of Chordoma
Jong-Beom Park, MD, Ulijinhu-H, South Korea; Choong-Ki Lee, MD, Seoul, South Korea

FDA DEVICE/DRUG STATUS: This abstract does not discuss or include any applicable devices or drugs.

5:31–5:56 p.m.
27. Effect of In Vivo Sustained Release of Growth Factors on Degenerated Disc Using a Rat Model
Scott Wengert, MD*; James Woodall, Jr, MD; Michelle Tudor, PhD; Ashley Nagah, MD; Ham Bengsted, PhD; Jackson, MS, USA

FDA DEVICE/DRUG STATUS: This abstract does not discuss or include any applicable devices or drugs.
2009 HTA TECHNOLOGY SELECTIONS

Background:
The Health Technology Assessment (HTA) program is a legislatively created program that seeks to ensure that health technologies purchased by state agencies are safe and effective, and that coverage decisions of state agencies are more consistent and transparent. The focus of the program is to rely on scientific, or evidence-based, information about safety and effectiveness to inform decisions and improve quality. An independent committee of eleven practicing health care clinicians uses the report and other information to review evidence regarding safety, efficacy, and cost-effectiveness of various medical procedures and/or equipment and determine if the state will pay for those procedures.

Technology Selection (Review or Re-review)
The HCA Administrator, in consultation with participating state agencies, selects technologies that undergo an evidence review, and may undergo a re-review. The independent committee can also be petitioned to include a technology for review or re-review, if not selected by the Administrator.

- State agency liaisons recommend potential technologies for prioritization and recommendation to the HCA Administrator. A priority ranking tool, with criteria based on legislative and other HTA program criteria, is used to rank the technologies.
- Interested individuals can petition the program to review a technology using the form located on the HTA website at: http://www.hta.hca.wa.gov/ipp.html. These petitions are also prioritized and a recommendation is made to the Administrator.
- Recommendations are posted for public and committee comment. The HCA Administrator considers the agency recommendation and public comment and selects technology topics. Topics are posted on our HTA website.
- Interested individuals may also petition the clinical committee to re-consider a topic or re-review of a technology that the HCA Administrator decided not to select.

Prioritization Criteria:
HTA created a process and tools based on the legislative requirements and criteria that are widely used in technology assessment priority settings. Identification of criteria and use of priority tools makes the process explicit and increases transparency and consistency across decision-makers. The tools are intended to be used by the medical directors when making recommendations and by the clinical committee when making comments or selections of technologies. The technology scoring tool has a corollary set of Primary criteria and a set of Secondary criteria. The primary criteria are directly linked to the legislative mandates for the program to focus technology reviews where there are concerns about safety, efficacy, or cost effectiveness, especially relative to existing alternatives. See RCW 70.14.100. These criteria are also common to other technology assessment programs.

Re-review Criteria
Re-review criteria are included in a prioritization document and directly linked to the legislative mandate that technologies shall be selected for re-review only where evidence has since become available that could change a previous determination. Technologies are considered for re-reviews at least once every eighteen months. Re-reviews consider only evidence made available since the previous determination. See RCW §70.14.100.

~ Based on these legislation, the re-review criterion is directed at identifying those situations:
- Where new evidence about a technology exists that was not available when the initial review was completed.
- And there is at least some likelihood that the new evidence could result in a change to a previous determination.
Factors here could include potentially high impact new outcomes not previously considered; new clinical trials that are high quality and have results that are counter to trials included in review; new cost or agency impact results.

Selection for 2009

- On September 23, 2008, the agency liaisons made recommendations on twelve potential technologies and two potential re-review technologies for prioritization.
- Topics and recommendations were posted to the website for comment.
- On December 12, 2008, the HCA Administrator selected seven topics for review and did not select any re-review topics. Selected topics are posted on our HTA website (http://www.hta.hca.wa.gov/).
- A stakeholder has requested that the clinical committee re-consider the Upright MRI as a topic for re-review.

2009 Re-review Recommendations:
The following re-review recommendations related to Upright MRI were made to the administrator.

| Upright MRI | A PubMed literature scan was completed by the HTA clinical consultant. It did not reveal any significant new evidence that might lead to a different conclusion. No other issues or concerns raised at this time, and this is not recommended for re-review by medical directors at this time. |

The Administrator also reviewed public comments submitted by stakeholders and after that review agreed with the initial recommendation that there was not sufficient new evidence that would likely lead to a different conclusion to warrant a re-review.
Health Technology Clinical Committee
Findings and Coverage Decision
Topic: Coronary Computed Tomographic Angiography
Meeting Date: November 14, 2008
Final Adoption:

Number and Coverage Topic
20081114A – Coronary Computed Tomographic Angiography for detection of Coronary Artery Disease.

HTCC Coverage Determination
Coronary Computed Tomographic Angiography (CCTA) is covered benefits with conditions consistent with the criteria identified in the reimbursement determination.

HTCC Reimbursement Determination

❖ Limitations of Coverage
1) Patients with low to intermediate risk of coronary artery disease;
2) For investigation of acute chest pain in an emergency department or hospital setting; and
3) Using Computed Tomography machines with 64-slice or better capability.

❖ Non-Covered Indicators
Patients who are asymptomatic or at high risk of coronary artery disease; CCTA used for coronary artery disease investigation outside of the emergency department or hospital setting; and CT scanners that use lower than 64-slice technology.

❖ Agency Contact Information

<table>
<thead>
<tr>
<th>Agency</th>
<th>Contact Phone Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Labor and Industries</td>
<td>1-800-547-8367</td>
</tr>
<tr>
<td>Uniform Medical Plan</td>
<td>1-800-762-6004</td>
</tr>
<tr>
<td>Health and Recovery Services Administration</td>
<td>1-800-562-3022</td>
</tr>
</tbody>
</table>
Computed Tomographic Angiography Background

The Computed Tomographic Angiography topic was selected and published in August 2007 to undergo an evidence review process. Heart disease is the leading cause of death and disability in the US: with 700,000 deaths. The most common heart disease in the US is coronary artery disease (CAD), which can lead to heart attack. CAD is a narrowing of one or more coronary arteries that result in an insufficient supply of oxygen to the heart muscle and is a leading cause of death in the US and developed countries. CAD may be asymptomatic or lead to chest pain (angina), heart attack, myocardial infarction (MI) or death. Non invasive tests include: Stress Echocardiograms – tests that compare blood flow with and without exercise and visualize the heart. Single-photon emission computed tomography (SPECT), also known as nuclear stress testing or myocardial perfusion imaging. Invasive tests include: The “gold” standard is the conventional coronary angiography which involves placement of a catheter and injection of contract material into a large artery or vein, followed by 2-dimensional visualization with x-rays. Coronary computed tomographic angiography (CCTA) is a minimally invasive radiological technique used to provide images of the heart and surrounding vessels.

CCTA has been suggested as an alternative or useful complementary approach to other non-invasive methods of diagnosing coronary artery disease (CAD). Due to its ability to visualize coronary anatomy, CCTA has been suggested as a strategy to rule out significant CAD among patients at low or intermediate risk of significant disease, thereby giving greater reassurance than other non-invasive methods and potentially reducing the number of patients ultimately sent for invasive coronary angiography (ICA). Potential drawbacks include radiation exposure; duplicative or additional testing; incidental findings; and uncertainty about whether the test results in better health outcomes.

In September 2008, the HTA posted a draft and then followed with a final report from a contracted research organization that reviewed publicly submitted information; searched, summarized, and evaluated trials, articles, and other evidence about the topic. The comprehensive, public and peer reviewed, Computed Tomographic Angiography report is 125 pages, identified 8 relevant studies for the Emergency room setting and 34 relevant studies for outpatient, Medicare coverage and 4 expert treatment guidelines. These studies represent the best available information; including a randomized controlled trial for the emergency room setting from which evidence based conclusions were drawn.

An independent group of eleven clinicians who practice medicine locally meet in public to decide whether state agencies should pay for the health technology based on whether the evidence report and other presented information shows it is safe, effective and has value. The committee met on November 14th, reviewed the report, including peer and public feedback, and heard public and agency comments. Meeting minutes detailing the discussion are available through the HTA program or online at http://www.hta.hca.wa.gov in the committee section.

Summary of Committee Findings

The committee found that it had the most complete information: a comprehensive and current evidence report, public comments, and agency utilization information. The committee concluded that the current evidence onComputed Tomographic Angiography demonstrates that there is sufficient evidence a decision about use in an emergency
setting to cover investigation of acute chest pain in an emergency room department or hospital setting for those who are at low-to-intermediate risk of coronary artery disease. The committee concluded that there is not sufficient, reliable evidence developed to make a determination for other coronary CTA uses, including the outpatient setting. For low-to-intermediate risk patients in the Emergency department setting the diagnostic accuracy of the 64-slice as a triage tool was supported by one RCT and several case series. For low-to-intermediate risk outpatients, no RCT or long-term cohort evidence was available. Modeling suggests a lower rate of false negatives than SECHO and SPECT, and a lower rate of false positives than SPECT, but these differences change with underlying prevalence of CAD and involves other trade-offs.

Based on these evidentiary findings, the committee voted: 2 for non-coverage and 7 for coverage with conditions.

- **Is it effective?**
The committee identified multiple key factors and health outcomes that were important for consideration in their overall decision on whether the technology was effective. Summary of committee consideration, discussion, and comments are listed below.

  ✓ **Diagnostic Accuracy – Sensitivity**: the committee agreed as a whole that CCTA has a high level of sensitivity. The technology report sensitivity rate was 98%; which compared favorably to stress echo at 76-94% and SPECT at 88-98%. The indeterminate rates were also lower, with CCTA at 3% versus Stress ECHO at 13% and SPECT at 9%.

  ✓ **Diagnostic Accuracy – Specificity**: the committee agreed equivalent specificity. Some uncertainty about lower prevalence population was shared amongst the committee members. The technology report specificity rate was comparable at 82-88%; compared to stress echo at 88% and SPECT at 77%.

  ✓ **Reduction in invasive CA**: the committee agreed that modeling suggests reduced ICA, but trial evidence data was inconclusive with Rubenstien trial showing reduction and Goldstein shiwoing slight increase, especially when compared to alternative diagnostic tools.

  ✓ **Replace other tests**: most modeled analysis and clinical trials used CCTA in conjunction with other tests. Committee agreed that CCTA wouldn’t replace other non-invasive technologies.

  ✓ **Incidental findings**: committee discussed as an issue both we respect to efficacy and safety and concluded that evidence demonstrates incidental findings are not infrequent events. Incidental findings can provide valuable information for diagnosis of previously undetected other diseases but also often leads to uncertainty or further tests to rule out questionable findings. The committee agreed that there is currently no evidence regarding improved patient health outcomes balancing cost and potential harms from further testing and anxiety.

  ✓ **Effect in real world**: Committee discussed several technology assessment key unknowns: whether more disease found will help or harm patients, especially at lower disease levels (clinical relevance is questionable); whether broad dissemination will result in lower test thresholds that may not result in better overall health outcomes but more radiation; and the extent to which CCTA can replace and not add to tests. Additionally, certification of machines and readers
was also discussed; hospitals require JAHCO accreditation and thus have some standards.

- **Is it safe?**
  The committee identified multiple key factors and health outcomes that were important for consideration in their overall decision on whether the technology was safe. Summary of committee consideration, discussion, and comments are listed below.

  ✓ **Radiation Exposure** is an important safety outcome to the committee. The committee discussed the technology assessment report findings of an overall cancer risk of .22% for women and .08% for men. Radiation dosage can be reduced through technique and machine type, but it is unknown whether these lowest dosage techniques/machines are used in WA settings. Overall exposure reported at between 2.0-8.0mSV for lower range is equivalent to SPECT; and 12.0 to 14.0 range for higher dose which is equivalent to A-bomb survivor at 2.3 kilometer distance. The committee concluded that there are small but definite risks, within appropriate norms. The radiation risks are high enough to obviate benefit when applied to very low risk patients.

  ✓ **Incidental findings** are also an important safety outcome that the committee discussed as an issue both we respect to efficacy and safety and concluded that evidence demonstrates incidental findings are not infrequent events. Incidental findings can provide valuable information for diagnosis of previously undetected other diseases but also often leads to uncertainty or further tests to rule out questionable findings. The committee agreed that there is currently no evidence regarding improved patient health outcomes balancing cost and potential harms from further testing and anxiety.

- **Does it provide value (improve health outcome)?**
  The committee discussed cost and cost-effectiveness as a whole. This topic generated the least discussion. There are several cost studies for ED and outpatient showing cost savings. The technology assessment report also modeled costs for ED and outpatient showing cost savings using Medicare reimbursement rates. No analysis included costs related to incidental findings or harms. Current state agency reimbursement rates do not correlate with modeled costs (Agency reimbursement for CCTA is higher and for comparators is lower).

  ✓ Committee members were split, with four considering the cost effectiveness currently unproven and five concluding that CCTA is either equivalent or more cost effective in some situations.

**Consistency with Medicare Decision and Expert Treatment Guidelines**

Committee reviewed and discussed the Medicare coverage decision and expert guidelines as identified and reported in the technology assessment report.

- There is no national coverage decision (NCD), however a coverage analysis and memo was issued in 2008 and summarized: there is uncertainty regarding any potential health benefits or patient management alterations from including coronary CTA in the diagnostic workup of patients who may have CAD. No adequately powered study has established that improved health outcomes can be casually attributed to coronary CTA for any well-defined clinical indication, and the body of evidence is of overall limited quality and limited applicability to Medicare patients.
with typical co-morbidities in community practice. The primary safety concerns with cardiac CTA are the exposure to radiation and the use of contrast and blocker medications.

- Four expert guidelines were identified that address the use of CCTA for detection of CAD, but not the setting (ED versus outpatient).
  - American Heart Association (2006): evidence supports the use of CCTA for patients with low-to-intermediate stenosis and may obviate the need for ICA.
  - Multi-Society Statement of Appropriateness Criteria for Cardiac Computed Tomography (2006): Appropriateness reviews deemed the use of CCTA for detection of CAD to be appropriate for the following patient populations: chest pain syndrome with intermediate pre-test probability of CAD and uninterpretable EKG or inability to exercise; chest pain and uninterpretable or equivocal stress test results; acute chest pain with intermediate pre-test probability of CAD and no EKG changes and serial enzymes negative; and symptomatic patients requiring evaluation of suspected coronary anomalies.
  - American College of Radiology (2006): CCTA is appropriate for assessment of CAD, although its usefulness for patients with low pre-test probability is unknown. Appropriateness rating of 7 out 9 for the evaluation of chronic chest pain.
  - SCCT/NASCI Consensus Update (2007): CCTA to be appropriate in the following circumstances: (1) to rule out significant coronary stenosis; (2) to evaluate patients with equivocal or discordant results on a stress perfusion or wall motion study; (3) to rule out stenosis in patients with a low pre-test likelihood of CAD and (4) to potentially replace diagnostic catheterization in patients undergoing non-coronary cardiac surgery.

The committee concluded that their decision is consistent with applicable policy and guidelines. There is no national Medicare coverage decision. The decision is consistent with treatment guidelines in that low to intermediate triage will be covered, with the coverage decision being more specific in identifying the place of service. The committee decision is based on all evidence, including public and agency comments and the comprehensive technology assessment report.

**Committee Authority**

Washington State believes it is important to use a scientific based, clinician centered approach for difficult and important health care benefit decisions. The HTA gathers and assesses the quality of the latest medical evidence using a scientific research company, takes public input at all stages, and asks a committee of eleven independent health care professionals to review all the information and render a decision at an open meeting. The Washington State Health Technology Clinical Committee (HTCC), an independent committee of 11 health practitioners, determines how selected health technologies are covered by several state agencies. See RCW 70.14.080-140. These technologies may include medical or surgical devices and procedures, medical equipment, and diagnostic tests. HTCC bases their decisions on the evidence of the technology’s safety, efficacy, and cost effectiveness. Participating state agencies are required to comply with the decisions of the HTCC. HTCC decisions may be re-reviewed at the determination of the HCA Administrator.
Missed Lumbar Disc Herniations Diagnosed With Kinetic Magnetic Resonance Imaging

Jun Zou, MD,* Huilin Yang, MD, PhD,* Masashi Miyazaki, MD,† Feng Wei, MD, PhD,† Soon W. Hong, MD,‡ Seung H. Yoon, MD,† Yuichiro Morishita, MD,† and Jeffrey C. Wang, MD†

Study Design. A novel dynamic magnetic resonance imaging (MRI) system, kinetic MRI (kMRI), was used to study lumbar disc herniations.

Objective. The objective of this study was to determine if adding flexion and extension MRI studies to the traditional neutral views would be beneficial in the diagnosis of lumbar disc herniations.

Summary of Background Data. Prior studies demonstrate that only 70% of patients with lumbar disc herniations based on physical examinations are confirmed by MRI studies. Recently, kMRI delivers the ability to scan patients in neutral, flexion, and extension positions, which may allow for improved diagnosis of this problem.

Methods. Five hundred fifty-three patients underwent kMRI with assessment of the degree of disc bulge in neutral and flexion and extension. The images were analyzed using computer measurement technology to objectively quantify the amount of disc herniation.

Results. For patients with normal or <3 mm of disc bulge in neutral, 19.46% demonstrated an increase in herniation to >3 mm bulge in extension, and 15.29% demonstrated an increase to >3 mm bulge in flexion. For patients in the neutral view that had a baseline disc bulge of 3 to 5 mm, 13.28% had increased herniations to >5 mm in extension and 8.47% had increased herniations to >5 mm in flexion. For patients with a baseline disc bulge of 5 to 7 mm in neutral, 10.58% increased in extension and 5.78% increased in flexion. In addition, for patients with a baseline disc bulge of 7 to 9 mm in neutral, 9.09% increased in extension and 4.55% increased in flexion.

Conclusion. A significant increase in the degree of lumbar disc herniation was found by examining flexion and extension views when compared with neutral views alone. kMRI views provide valuable added information, especially in situations where symptomatic radioluclarity is present without any abnormalities demonstrated on conventional MRI.

Key words: kinetic MRI, lumbar disc herniation, missed diagnosis. Spine 2008;33:E140–E144

Lower back pain is the second most common reason for physician visits in the United States, second only to colds and flu.1 Americans spend at least $50 billion each year on low back pain. One common reason for lower back pain is herniation of the intervertebral disc into the spinal canal. In the United States in 2003, the National Hospital Discharge Survey reported that 3,57,000 procedures were performed for disorders of the intervertebral disc,2 8.5% higher than in 2000.3

Magnetic resonance imaging (MRI) is a good tool most frequently used for lumbar disc herniation because it can show abnormal areas of soft tissue around the spine. However, traditional MRI has significant limitations, although it reveals musculoskeletal disease. The patients are placed in a horizontal, non-weight-bearing position where conventional scans may not reveal the causative pathology. However, only 70% of patients who were diagnosed with a lumbar disc herniation based on clinical examination had a lumbar disc herniation confirmed by MRI.4

Recently, kinetic MRI (kMRI) permits us to image the patient in a weight-bearing position (either standing up or sitting), and in the flexed and extended positions, which can, of course, reveal abnormalities that were missed by a conventional MRI study. It may supply a more thorough investigation of each patient and allow us to better understand the true nature of the pathology. Imaging the spine in the weight-bearing position with extension and flexion or placing the spine in the position of pain may increase the diagnostic accuracy for the surgeons. The purpose of this study was to study the use of kMRI for evaluation of missed herniated discs when compared with conventional MRI studies and to determine the changes in the disc herniations according to the flexed and extended positions.

Materials and Methods

Study Population

From July 2005 through July 2006, 553 patients with symptomatic back pain with/without radiculopathy were referred to kMRI for lumbar MRI examination. There were 234 males and 319 females. The mean age was 46.2 years (range, 18–76 years). This represented 2765 lumbar discs in total.

Imaging Instrumentation

MRI of the lumbar spine was performed by using a 0.6 Tesla MRI scanner (Fonar Corp. UPRIGHTTM, Multi-Position, NY, NY). The MR unit uses a vertical orientation of the 2 opposing magnet doughnuts, allowing scanning of the patient in an upright axially loaded position. An 18-inch gap between the magnets is present. Images were obtained using a quad channel planar coil. T1 weighted sagittal spin echo images (repetition time 671 milliseconds, echo time 17 mil-
Dynamic Change in Lumbar Disc Herniations During Lumbar Extension and Flexion

On extension images, the pair T-test showed significant increases in disc herniation from the neutral position to the extension position at each level (P < 0.005). The results were as followed: L1/L2 (2.12 ± 1.06 vs. 2.39 ± 1.83 mm), L2/L3 (2.44 ± 1.24 vs. 2.69 ± 1.77 mm), L3/L4 (2.78 ± 1.28 vs. 3.08 ± 2.25 mm), L4/L5 (3.48 ± 1.59 vs. 3.82 ± 2.47 mm), and L5/S1 (3.45 ± 1.78 vs. 3.77 ± 2.58 mm). On extension images, the pair T-test showed significant differences only at L3/L4 and L4/L5 from the neutral position to the flexion position, L3/L4 (2.78 ± 1.28 vs. 2.68 ± 1.33 mm), L4/L5 (3.48 ± 1.59 vs. 3.34 ± 1.57 mm) (P < 0.05). There were no significant changes at L1/L2 (2.12 ± 1.06 vs. 2.15 ± 1.24 mm), L2/L3 (2.44 ± 1.24 vs. 2.36 ± 1.30 mm), and L5/S1 (3.45 ± 1.78 vs. 3.34 ± 1.70 mm) (P > 0.05) (Figure 1).

Table 1. Distribution of Lumbar Disc Herniation Among Neutral, Extension, and Flexion Images

<table>
<thead>
<tr>
<th>Grade</th>
<th>Grade 1 (0–3 mm)</th>
<th>Grade 2 (3–5 mm)</th>
<th>Grade 3 (5–7 mm)</th>
<th>Grade 4 (7–9 mm)</th>
<th>Grade 5 (&gt;9 mm)</th>
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<tr>
<td>Neutral</td>
<td>1557</td>
<td>956</td>
<td>208</td>
<td>44</td>
<td>0</td>
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<tr>
<td>Extension</td>
<td>1254</td>
<td>1132</td>
<td>313</td>
<td>62</td>
<td>4</td>
</tr>
<tr>
<td>Flexion</td>
<td>1319</td>
<td>1113</td>
<td>277</td>
<td>54</td>
<td>2</td>
</tr>
</tbody>
</table>

Figure 1. Bar graph shows dynamic changes in lumbar disc herniation during lumbar extension and flexion. The values of lumbar disc herniation mean ± SD in the extension and flexion views were compared with the neutral view (*P < 0.05, **P < 0.01).
1319 (84.71%) of 1557 discs and progressed to grade 2 in 238 (15.29%). In the grade 2 group, disc bulge in 875 (91.53%) of the 956 discs maintained grade 2 and progressed to grade 3 in 81 (8.47%). In the grade 3 group on extension MRI, disc bulge was more severe than that in neutral position in 12 (5.78%) of the 208 discs and was maintained at grade 3 in 196 (94.22%). In the grade 4 group, disc bulge in 42 (95.45%) of the 44 discs maintained grade 4 and progressed to grade 5 in 2 (4.55%) (Figure 2). The Fisher’s exact test showed significant difference these 4 groups for increasing disc herniation during flexion \( (P < 0.05) \). With regard to the grade of disc herniation, the \( \chi^2 \) test was used to examine the difference between extension and flexion imaging. There are significant differences in all grades \( (\chi^2 = 16.19, 14.11, 5.06, \text{respectively, } P < 0.05) \) except grade 4 \( (\chi^2 = 0.5, P > 0.05) \) (Table 2).

**Table 2. Incidence of Missed Diagnosis of Lumbar Disc Herniation Showed by Extension and Flexion Images**

<table>
<thead>
<tr>
<th>Grade of Disc Bulge (in the Neutral Position)</th>
<th>Extension</th>
<th>Flexion</th>
<th>( \chi^2 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grade 1 (( \leq 3 ) mm)</td>
<td>19.46% (303/1557)</td>
<td>15.29% (238/1557)</td>
<td>16.19*</td>
</tr>
<tr>
<td>Grade 2 (3–5 mm)</td>
<td>13.28% (127/956)</td>
<td>8.47% (81/966)</td>
<td>14.11*</td>
</tr>
<tr>
<td>Grade 3 (5–7 mm)</td>
<td>10.58% (122/208)</td>
<td>5.78% (12/208)</td>
<td>5.06†</td>
</tr>
<tr>
<td>Grade 4 (7–9 mm)</td>
<td>9.09% (4/44)</td>
<td>4.55% (2/44)</td>
<td>0.5</td>
</tr>
</tbody>
</table>

* \( P < 0.01 \).
† \( P < 0.05 \).

Since chronic low back pain is not a single factorial disease, it has various etiologies. However, low back pain has been believed in some cases to be related to the intervertebral disc, the surrounding soft tissue, or the facet joints. MRI has become the examination of choice for diagnosing lumbar disc herniation.\(^5\) The pros of it is that it has no known side effects, no radiation exposure, and noninvasive. In fact, weight bearing, flexion, extension, or lateral bending may change anatomic relationships. Compressive load can increase the load in the lumbar spine by 80% compared with that in the supine position.\(^6\) In addition, the intradiscal pressure also changes with the position of spine where it increases in standing, sitting, and in a forward flexed position.\(^7\) Prolonged standing can diminish the size of the neural foramens and central spinal canal because the discs lose water content and height whenever the load on the spine is increased. Axial loading of the spine decreases the disc height measured on MR images, and axial compression of the spine also causes bulging of the intervertebral disc and narrowing of the diameters of the neural foramens and central canal. Scanning patients in a recumbent position may potentially miss an occult disc herniation, which may be revealed in a weight-bearing or more positional mode such as flexion or extension. Radiologists failed to report certain pathologic findings, which had to be handled during the surgery. Cases where there is such limited association between diagnostic imaging and clinical symptoms perplexed surgeons a long time. Flexion and extension radiographs and computed tomography myelography were the standard methods of obtaining positional images of the spine. However, because MRI yields an image that is superior to radiographs and less invasive than myelography, physicians have been experimenting with ways of using MRI to obtain positional images of spine. To help in a better understanding of the pathophysiology of the spine, there seems to be a need for further developments in functional clinical imaging.

Cartolari\(^8\) set up an axial-loaded computed tomography and MR technology by pressing on the recumbent patients’ shoulders with 70% body weight. Smith\(^9\) figured out a study of 25 patients with low back pain and sciatica for lumbar spine upright MRI. Upright MRI demonstrated abnormalities in 13 patients (52%) that were not evident in the recumbent posture. There were 3 cases with lateral disc herniations, 6 cases with hypermo-
altered mobility is related to symptoms. Karadimas et al. healthy spines, which is important of understanding how movement technique influenced segmental motion of an increased lumbar lordosis. That indicted how passive Force applied on the other vertebrae (L3–L5) resulted in (L1–L2) consequently decreased the lumbar lordosis. Anterior force applied at the upper lumbar vertebrae directed force was applied manually at each lumbar spinotic subjects were positioned prone within a vertically mobile of the lumbar spine during a posterior to anterior to 15.29% (238/1557). This also suggests that extension MRI, the incidence of missed disc herniations is up to 19.46% (303/1557). Using only flexion MRI compared with conventional MRI, the incidence of missed disc herniations is up to 16.49% vs. 333 discs, 12.04%). Flexion and extension MRI views provide valuable, added information when assessing patients for lumbar disc herniations, and may be especially useful in situations where symptomatic radiculopathy is present with unimpressive conventional MRI studies.

This imaging technology may prove to be useful to reveal hidden pathologies not only in occult disc herniations, but also in the other degenerative spinal disease. kMRI may be able to detect occult stenosis or occult instability in the spine by placing the patient’s spine in the position that causes pain or in a position that should narrow the spinal canal and neural foramen (such as spinal extension). In addition, large or claustrophobic patients or patients who need to be scanned in an upright position because of congestive heart failure, severe chronic obstructive pulmonary disease, or severe spinal kyphosis, can be handled by this novel MRI.

### Key Points
- Prior studies demonstrate that only 70% of patients who were clinically diagnosed with lumbar disc herniations based on physical examinations had lumbar disc herniations confirmed by MRI studies.
- A novel dynamic magnetic resonance imaging system, Kinetic MRI (kMRI), delivers the ability to scan patients in an upright position that causes pain or in a position that should narrow the spinal canal and neural foramen (such as spinal extension). In addition, large or claustrophobic patients or patients who need to be scanned in an upright position because of congestive heart failure, severe chronic obstructive pulmonary disease, or severe spinal kyphosis, can be handled by this novel MRI.

### References
Kinematic Analysis of the Relationship Between the Grade of Disc Degeneration and Motion Unit of the Cervical Spine

Masashi Miyazaki, MD,* Soon Woo Hong, MD, PhD,* Seung Hwan Yoon, MD, PhD,* Jun Zou, MD,* Benjamin Tow, MD,* Ahmet Alanay, MD,† Jean-Jacques Abitbol, MD,‡ and Jeffrey C. Wang, MD*

**Study Design.** Kinetic MRIs of cervical spines were obtained and analyzed according to the amount of motion and the degenerative grade of the intervertebral disc.

**Objective.** To define the relationship between the grade of disc degeneration and the motion unit of the cervical spine and elucidate changes in the role of each cervical spine unit during flexion-extension motion caused by degeneration.

**Summary of Background Data.** Degenerative changes in the cervical disc occur with age. The correlation between the degree of cervical disc degeneration and extent of cervical spine mobility has not yet been determined. The effect of degeneration on the overall motion of the functional spinal unit also remains undefined.

**Methods.** We studied 164 patients with symptomatic neck pain. The cervical intervertebral discs were graded by spine surgeons according to the degenerative grading system (Grades I to V). All radiologic data from kinetic MRIs were recorded on a computer for subsequent measurements. All measurements and calculations for translational motion and angular variation of each segment were automatically performed by a computer analyzer.

**Results.** The translational motion in discs with Grade II degeneration (mild degeneration) increased to Grade III degeneration (higher degeneration). However, the translational motion and angular variation significantly decreased for the Grade V (severe degeneration). For patients with relatively low grades of degeneration, Grades I and II discs, the C4–C5 and C5–C6 segmental units contributed the majority of total angular mobility of the spine. However, for the severely degenerated segments, Grade V discs, the contributions of the C4–C5 and C5–C6 U significantly decreased.

**Conclusion.** The changes that occur with disc degeneration progress from the normal state to an unstable phase with higher mobility and subsequently to an ankylosed stage. This study evaluated the contribution of different levels to the changes in overall motion that occur with degeneration.

**Key words:** kinematic analysis, disc degeneration, cervical spine, intervertebral disc, kinetic magnetic resonance imaging. *Spine* 2008;33:187–193

Cervical disc degeneration is common after middle age and is part of the normal aging process.¹⁻⁴ The degenerative changes generally occur gradually and frequently without significant symptoms; however, some individuals become symptomatic. Progressive degeneration affects the motion units and overall kinematics of the cervical spine.⁵ One of the primary functions of the cervical spine is to control head and neck motion. Cervical instability has long been considered as a contributor to neck and back pain.⁶⁻⁹ Many motion studies have been performed to define normal flexibility and to determine the relationship between abnormal motion and neck and related pain conditions.¹⁰⁻¹³ The correlation between the degree of cervical disc degeneration and the extent of cervical spine mobility has not yet been determined. The effect of degeneration on the overall motion of the functional spinal unit also remains undefined.

Many investigators have measured this motion using simple flexion and extension radiographs.¹⁰,¹²,¹³ The accuracy of an analysis performed using simple radiographs depends on methodology. Generally, it increases with the complexity of the methodology; however, its practicality in routine clinical applications consequently decreases. The radiostereometric method provides accurate tracking of motion¹⁴; however, it necessarily involves the implantation of metal markers. Hence, the use of this invasive method is not practical in many clinical scenarios.

Kinetic MRI studies can noninvasively demonstrate the mobility of each motion segment and define the region of spin motion.¹⁵⁻¹⁷ Degeneration is caused by the loss of both proteoglycan and water in the disc and is detected by the loss of signal intensity on T2-weighted images.¹⁸ It induces structural changes in the disc and a decrease in disc height, which can be visualized on MR images.¹⁹,²⁰ Thus, kinetic MRI can relate the grade of degeneration to the extent of cervical spine mobility. In addition, a recently developed computer-assisted calculation method facilitates the measurement of segmental...
angular variation and translational motion and the comparison of different positional (flexion/neutral/extension) MRIs.\textsuperscript{10,11}

In this study, the relationship was studied between the grade of disc degeneration and the motion of the cervical spine units in patients with cervical pain using kinetic MRI. In addition, this study aimed to elucidate changes in the role of each cervical spine unit during flexion-extension motion in patients with disc degeneration using kinetic MRI.

\section*{Materials and Methods}

\subsection*{A Grading System for Cervical Intervertebral Disc Degeneration and Its Reliability}

A comprehensive grading system for cervical disc degeneration was developed based on the reported literature\textsuperscript{21–25} (Table 1, Figure 1). The cervical intervertebral discs were graded by spinal surgeon observers according to this grading system by using T2-weighted sagittal images.

\begin{table}[h]
\centering
\begin{tabular}{|c|c|c|c|}
\hline
Grade & Nucleus Signal Intensity & Nucleus Structure & Distinction of Nucleus and Annulus & Disc Height \\
\hline
I & Hyperintense & Homogenous, white & Clear & Normal \\
II & Hyperintense & Inhomogenous with horizontal band, white & Clear & Normal \\
III & Intermediate & Inhomogenous, gray to black & Unclear & Normal to decreased \\
IV & Hypointense & Inhomogenous, gray to black & Lost & Normal to decreased \\
V & Hypointense & Inhomogenous, gray to black & Lost & Collapsed \\
\hline
\end{tabular}
\caption{Grading System for Cervical Intervertebral Disc Degeneration}
\end{table}

\subsection*{Intra/Interobserver Reliability.}

T2-weighted sagittal images were used by 3 spine surgeon observers (A–C) to Grade 300 cases of cervical intervertebral disc degeneration in a blinded fashion. All the MRIs were separately analyzed with a minimum interval of 1 week. A set of typical sample MRIs (Table 1, Figure 1) was available to the observers during the image review.

\subsection*{Participants}

Kinetic MRI scans of the cervical spine were consecutively obtained over a 6-month period from February 2006 to July 2006. In this study, 168 patients with symptomatic neck pain with/without radiculopathy or myelopathy were enrolled; of these, 164 consecutive patients (69 men and 95 women) were selected based on the image findings. The mean age of the selected patients was 44 years (range, 19–93 years). All the selected

![Figure 1. Grading system for cervical intervertebral disc degeneration. A, Grade I: Nucleus signal intensity is hyperintense and nucleus structure is homogeneous, white. Distinction of nucleus and anulus is clear. Disc height is normal. B, Grade II: Nucleus signal intensity is hyperintense and nucleus structure is inhomogeneous with horizontal band, white. Distinction of nucleus and annulus is clear. Disc height is normal. C, Grade III: Nucleus signal intensity is intermediate and nucleus structure is inhomogeneous, gray to black. Distinction of nucleus and annulus is unclear. Disc height is normal to decrease. D, Grade IV: Nucleus signal intensity is hypointense and nucleus structure is inhomogeneous, gray to black. Distinction of nucleus and annulus is lost. Disc height is normal to decrease. E, Grade V: Nucleus signal intensity is hypointense and nucleus structure is inhomogeneous, gray to black. Distinction of nucleus and annulus is lost. Disc height is collapsed. Grading was performed on T2-weighted midsagittal images.](image1)

![Figure 2. In each film, 77 points were marked for digitization.](image2)
patients were referred for cervical kinetic MRI scans because they exhibited cervicobrachial pain symptoms. None of the patients had previously undergone cervical spinal surgery. This study selected 6 level units (C2–C3, C3–C4, C4–C5, C5–C6, C6–C7, and C7–T1) and assessed a total of 492 T2-weighted midsagittal images.

**MRI Positioning**

With each patient in a sitting posture, neck flexion and extension were actively performed. The same amount of flexion extension is performed in all the 3 positions (40° flexion, 0° neutral, and −20° extension positions).

**Image Analysis**

MR imaging of the cervical spine was performed using a 0.6-Tesla MRI scanner (Fonar Corp. Upright Multi-Position, New York, NY) and a flexible surface coil. The MR unit uses a vertical orientation of 2 opposing magnet doughnuts, permitting scanning of the patient in an upright axially loaded position. There was an 18-in gap between the magnets. The patients were examined using T1-weighted sagittal spin-echo images (repetition time, 671 ms; echo time, 17 ms; thickness, 3.0 mm; field of view, 24 cm; matrix, 256 × 200; number of excitations (NEX), 2) and T2-weighted sagittal fast spin-echo images (repetition time, 3432 ms; echo time, 160 ms; thickness, 3.0 mm; field of view, 24 cm; matrix, 256 × 224; NEX, 2).

All radiologic data obtained from the MRIs were recorded on a computer for subsequent measurements, and all calculations were automatically performed on true MRI by a MRI analyzer. For each image, 77 points were marked for digitization by the spine surgeons. Specific points were selected for the occiput (Oc) and C1 and C2 vertebrae. Anterior and posterior baselines were marked at the Oc. Anterior tubercle and posterior margin of the atlas and the lowest end of the spinous process were marked at C1. At C2, 1 point was marked at the tip of the odontoid process and others, corresponding to C3 to T1. For each of the typical cervical vertebrae from C3 to T1, the vertebral body was denoted by 4 points (anterior-inferior, anterior-superior, posterior-superior, and posterior-inferior corners); disc height, by 2 points (middle of endplate); and the pedicle and spinal cord diameters, by 2 points each (Figure 2).

The basic measurements involved calculations of all the static intervertebral angular displacements and translations in different positions. Subsequently, total flexibility (motion segment integrity, translational motion, and angular variation) was calculated at each vertebral level from the difference in flexibilities in the flexion and extension positions. Translational motion was measured in 6 levels (C2–C3, C3–C4, C4–C5, C5–C6, C6–C7, and C7–T1 level) by determining the anterioposterior motion of 1 vertebra over another; a positive value indicated anterior translation (antelisthesis), whereas a negative value indicated posterior translation (retrolisthesis). Angular variation was measured in 5 levels (C2–C3, C3–C4, C4–C5, C5–C6, and C6–C7 level). For determining angular variation, lines were drawn through the inferior borders of 2 vertebral bodies adjacent to the corresponding vertebral levels. The lordotic angle was defined as negative, whereas the kyphotic angle, as positive (Figure 3). Both translational motion and angular variation, total movements (flexion to extension) were calculated at each level. Total number of assessed discs for translation motion was 984 discs, and total number of assessed discs for angular variation was 820 discs.

To elucidate changes in the role of each cervical spine unit during flexion-extension motion caused by degeneration, we estimated the contribution of each intervertebral level to total angular mobility.

Contribution of each level to total angular mobility (%) = (angular movement of each unit in degrees)/(angular movement of C1–C2 + C2–C3 + C3–C4 + C4–C5 + C5–C6 + C6–C7 in degrees) × 100.

**Statistical Analysis**

Statistical analysis was performed using the computer program SPSS (version 13, SPSS Inc., Chicago, IL), and values were expressed as mean ± standard deviation (SD). Student t test was performed with a significance level of 0.05. Intra and interobserver Kappa reliability was evaluated (Table 3).

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**Table 2. Cervical Disc Degeneration Grading**

<table>
<thead>
<tr>
<th>Level</th>
<th>Grade I</th>
<th>Grade II</th>
<th>Grade III</th>
<th>Grade IV</th>
<th>Grade V</th>
</tr>
</thead>
<tbody>
<tr>
<td>C2–C3</td>
<td>12</td>
<td>19</td>
<td>75</td>
<td>58</td>
<td>0</td>
</tr>
<tr>
<td>C3–C4</td>
<td>12</td>
<td>18</td>
<td>66</td>
<td>60</td>
<td>8</td>
</tr>
<tr>
<td>C4–C5</td>
<td>9</td>
<td>12</td>
<td>73</td>
<td>53</td>
<td>17</td>
</tr>
<tr>
<td>C5–C6</td>
<td>8</td>
<td>16</td>
<td>57</td>
<td>40</td>
<td>42</td>
</tr>
<tr>
<td>C6–C7</td>
<td>17</td>
<td>35</td>
<td>57</td>
<td>29</td>
<td>26</td>
</tr>
<tr>
<td>C7–T1</td>
<td>33</td>
<td>60</td>
<td>55</td>
<td>10</td>
<td>6</td>
</tr>
</tbody>
</table>

**Table 3. Intra- and Interobserver Reliability**

<table>
<thead>
<tr>
<th>Intraobserver</th>
<th>Kappa</th>
<th>Interobserver</th>
<th>Kappa</th>
</tr>
</thead>
<tbody>
<tr>
<td>A1–2</td>
<td>0.907</td>
<td>A1–B1</td>
<td>0.779</td>
</tr>
<tr>
<td>B1–2</td>
<td>0.950</td>
<td>A1–C1</td>
<td>0.752</td>
</tr>
<tr>
<td>C1–2</td>
<td>0.933</td>
<td>B1–C1</td>
<td>0.720</td>
</tr>
</tbody>
</table>
The interobserver reliability of the MRI evaluations was estimated using kappa statistics and interpreted according to the guidelines suggested by Landis and Koch. The levels of agreement were rated as follows: kappa value of 0 to 0.2 indicated poor agreement; 0.21 to 0.4, fair agreement; 0.41 to 0.60, moderate agreement; 0.61 to 0.8, substantial agreement; and 0.81, upward excellent agreement. A value of 1 indicated absolute agreement; whereas 0, agreement no better than chance. A kappa value of 0 to 0.2 indicated poor agreement; 0.21 to 0.4, fair agreement; 0.41 to 0.60, moderate agreement; 0.61 to 0.8, substantial agreement; and 0.81, upward excellent agreement. A value of 1 indicated absolute agreement; whereas 0, agreement no better than chance.

### Results

#### Grades of Cervical Disc Degeneration in the Study Population

A total of 984 cervical discs were graded in a study population of 164 individuals. The amount of disc degeneration corresponding to each grade is summarized in Table 2. For Grade I (almost normal) discs, the amount of disc degeneration is relatively lower at the C4–C5 and C5–C6 levels than at the other levels. In contrast, it is higher at the C4–C5, C5–C6, and C6–C7 levels than at the C2–C3, C3–C4, and C7–T1 levels for Grade V (severe degeneration) discs.

#### Kappa Values for Intra- and Interobserver Agreement

The estimated kappa values for intra and interobserver agreement are summarized in Table 3. Intraobserver agreement among the 3 readers was “excellent” with kappa values ranging from 0.907 to 0.950. Interobserver agreement was expected to be relatively lower than intraobserver agreement; however, the former was found to be “substantial” with kappa values ranging from 0.730 to 0.779.

#### Translational Motion

The mean values of translational motion for each cervical unit are shown in Table 4, and total intervertebral translational motion is shown in Figure 4. On each cervical unit, with increasing degenerative grade at same level, translational motion in Grade II (mild degeneration) tended to increase for segments with discs with Grade III (higher degeneration). However, the translational motion of the segments was observed to significantly decrease in Grade V discs. Significant differences in translational motion were observed between Grades II and III discs at the C7–T1 level and in the total intervertebral translational motion, and between Grades IV and V discs at the C4–C5 and C5–C6 levels and also in the total intervertebral translational motion.

#### Angular Variation

The mean values of angular variation for each cervical unit are shown in Table 5 and total intervertebral angular motion is shown in Figure 5. No significant difference was observed in the angular variation of each cervical unit in Grades I to IV discs relative to their immediate less degenerative grade at the same level. However, the angular variation of the segments was observed to significantly decrease in Grade V discs. Significant difference in angular variation was observed between Grades IV and V discs at the C4–C5 and C5–C6 levels and in the total intervertebral angular variation.

#### Contribution of Each Intervertebral Level to Total Angular Mobility

The contribution of each intervertebral level to total angular mobility is shown in Table 6. In Grades I and II discs, the role of C4–C5 and C5–C6 segmental units contributed the majority of total angular mobility of the spine. For Grades III and IV, the motion in segments C3–C4 and C6–C7 increased, as did the motion in segments C4–C5 and C5–C6. There was significant difference on Grades III, IV, and V compared to Grade I at C3–C4 level. In Grade V discs, the contribution of the C4–C5 and C5–C6 U significantly decreased. Significant

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Table 4. Translational Motion for Each Cervical Unit (mm)

<table>
<thead>
<tr>
<th>Level</th>
<th>Grade I</th>
<th>Grade II</th>
<th>Grade III</th>
<th>Grade IV</th>
<th>Grade V</th>
</tr>
</thead>
<tbody>
<tr>
<td>C2–C3</td>
<td>1.02 ± 0.90</td>
<td>0.96 ± 0.79</td>
<td>1.12 ± 0.75</td>
<td>1.17 ± 0.92</td>
<td>NA</td>
</tr>
<tr>
<td>C3–C4</td>
<td>1.36 ± 0.78</td>
<td>1.16 ± 0.96</td>
<td>1.40 ± 0.83</td>
<td>1.36 ± 0.84</td>
<td>0.95 ± 0.66</td>
</tr>
<tr>
<td>C4–C5</td>
<td>1.19 ± 0.71</td>
<td>1.55 ± 0.73</td>
<td>1.22 ± 0.91</td>
<td>1.32 ± 0.89</td>
<td>0.88 ± 0.59*</td>
</tr>
<tr>
<td>C5–C6</td>
<td>0.71 ± 0.53</td>
<td>1.15 ± 0.80</td>
<td>1.16 ± 0.80</td>
<td>1.42 ± 1.38</td>
<td>0.90 ± 0.70*</td>
</tr>
<tr>
<td>C6–C7</td>
<td>0.82 ± 0.68</td>
<td>0.88 ± 0.56</td>
<td>1.08 ± 1.37</td>
<td>0.85 ± 0.69</td>
<td>1.01 ± 0.80</td>
</tr>
<tr>
<td>C7–T1</td>
<td>0.69 ± 0.46</td>
<td>0.64 ± 0.50</td>
<td>1.01 ± 0.64*</td>
<td>1.00 ± 0.70</td>
<td>0.63 ± 0.40</td>
</tr>
</tbody>
</table>

*Values are expressed mean ± SD.

$p < 0.05$ compared to immediate less degenerative grade at same level.
difference was noted in the contribution of the C4–C5 and C5–C6 levels in Grades V and I discs.

**Discussion**

Intervertebral disc degeneration is an age-related physiologic process caused by a decrease in the proteoglycan and water concentration in the disc. MRI is the most accurate method to evaluate intervertebral disc morphology and to diagnose disc degeneration. Cervicobrachial pain may originate from cervical disc degeneration, which is associated with abnormal instability. However, the natural history of the change in cervical spine motion units following degeneration has not been sufficiently elucidated.

The pathomechanics of a degenerative lumbar spine has been described by Kirkaldy-Willis and Farfan. They postulated 3 stages of degeneration with different stability and motion conditions. The first stage was a temporary dysfunction phase characterized by early signs of disc degeneration and fibrillation of the articular cartilage of the facet joints. The second stage was the unstable phase characterized by abnormal movement of the spinal units. A decrease in proteoglycan and water concentration in the intervertebral disc weakens the structure, thereby restricting movement between adjacent spinal units. The disc height reduces, facet capsule loosens, and finally, the joint subluxates. In the last stage, the deformity is assumed to be stabilized by osteoligamentary repair mechanisms. At an advanced stage of degeneration, secondary responses occur in the adjacent osseous and soft tissue structures, causing restabilization of the spine.

Using a cadaver spine, Tanaka et al demonstrated that the kinematic properties of the lumbar spine are related to disc degeneration. They graded the severity of disc degeneration using MRI and cryomicrotome sections and measured the motion segments under multidirectional loading conditions. Their results demonstrated that greater motion was observed in the higher degeneration stage, i.e., in discs with radial tears of the annulus fibrosus. The disc space collapsed as found in severe degeneration stage resulted in stabilization of the motion units. Axelsson et al performed a radiostereometric analysis to investigate changes in intervertebral mobility caused by degeneration. They graded disc degeneration based on conventional radiography and MRI findings. Although they were unable to verify the unstable phase, they concluded that the stabilization stage was apparent when the disc height decreased by more than 50%. Dai investigated the correlation between disc degeneration and cervical instability. He graded disc degeneration according to T2 signal intensity observed on MRI and measured horizontal and angular instability using conventional lateral flexion and extension radiography. Cervical segmental instability was correlated with a higher degeneration stage.

In this study, we graded cervical disc degeneration using MRI and measured the motion with kinetic MRI. We have demonstrated the changes in translational motion and angular variation for each cervical unit following degeneration. Translational motion changed as the disc degenerated from its normal state to an unstable phase and finally to an ankylosed stage with increased stability. Angular variation significantly decreased in severe degeneration. Although our study focused on the cervical spine, our results support the hypothesis proposed by Kirkaldy-Willis and Farfan. We believe that kinetic MRI can be applied to clinical practice for defining spinal intervertebral mobility and for determining the treatment mode by studying the progression of disc degeneration, i.e., whether the disc eventually stabilizes or continues to degenerate.

**Table 5. Angular Variation for Each Cervical Unit (Degree)**

<table>
<thead>
<tr>
<th>Level</th>
<th>Grade I</th>
<th>Grade II</th>
<th>Grade III</th>
<th>Grade IV</th>
<th>Grade V</th>
</tr>
</thead>
<tbody>
<tr>
<td>C2–C3</td>
<td>6.94 ± 4.72</td>
<td>5.42 ± 4.46</td>
<td>6.94 ± 5.02</td>
<td>6.05 ± 4.27</td>
<td>NA</td>
</tr>
<tr>
<td>C3–C4</td>
<td>7.74 ± 9.86</td>
<td>8.02 ± 5.28</td>
<td>10.51 ± 6.07</td>
<td>9.87 ± 5.35</td>
<td>9.04 ± 5.28</td>
</tr>
<tr>
<td>C4–C5</td>
<td>11.54 ± 2.54</td>
<td>11.76 ± 7.43</td>
<td>10.07 ± 6.88</td>
<td>10.56 ± 6.65</td>
<td>6.62 ± 4.53*</td>
</tr>
<tr>
<td>C5–C6</td>
<td>12.06 ± 7.52</td>
<td>9.59 ± 6.94</td>
<td>10.51 ± 6.16</td>
<td>10.01 ± 7.87</td>
<td>6.27 ± 4.10*</td>
</tr>
<tr>
<td>C6–C7</td>
<td>8.51 ± 5.36</td>
<td>9.76 ± 5.38</td>
<td>8.15 ± 3.30</td>
<td>7.68 ± 5.46</td>
<td>7.59 ± 5.07</td>
</tr>
</tbody>
</table>

Values are expressed mean ± SD.

*P < 0.05 compared to immediate less degenerative grade at same level.

![Figure 5](image_url)
According to previous studies, the most commonly affected level is the intervertebral disc at C5–C6 followed by C6–C7 and C4–C5. The C2–C3 level is the least commonly affected. In our retrospective investigation on 164 patients, we observed that the amount of disc degeneration at the C4–C5 and C5–C6 levels was less than that at the other levels in normal discs. In contrast, it was greater at the C4–C5, C5–C6, and C6–C7 levels than at the C2–C3, C3–C4, and C7–T1 levels in severe degeneration. These results prompted us to assume that the C4–C5, C5–C6, and C6–C7 levels contributed the majority to the overall motion of the cervical spine from the early degenerative stage. Using dynamic radiography, Lind et al. evaluated the normal range of motion of the cervical spine in healthy subjects and observed that the greatest proportion of intersegmental flexion-extension motion occurred between the C4–C5 and C5–C6 levels and that a linear decrease in motion occurred with age. Using lateral dynamic radiography, Holmes et al. measured angular motion ranges from C2 to C7 in normal subjects and in those with cervical myelopathy and obtained findings similar to those reported by Lind et al. Using conventional lateral dynamic radiography in asymptomatic subjects, Lin et al. demonstrated that the greatest sagittal motion occurred between C4–C5 and C5–C6 and the least motion, at C2–C3. Most studies used active motion for calculating segmental angulation and obtained consistent data. However, these studies did not take into consideration the change in motion that occurs with degeneration. Therefore, we investigated the change in the role of each cervical spine unit during flexion-extension motion caused by degeneration. Our results indicated that the C4–C5 and C5–C6 segmental units contributed the majority of total angular mobility of the spine in normal discs. The contribution of the C3–C4, C4–C5, C5–C6, and C6–C7 U increased in advanced degeneration. The roles of the C4–C5 and C5–C6 U in total angular mobility significantly decreased in severe degeneration.

From these results, we hypothesized that cervical disc degeneration begins at the C4–C5 and C5–C6 levels because these units withstand the maximum load for the overall motion in the early phase. Following degeneration, these segmental units undergo changes from an unstable phase to an ankylosed stage. Many clinical studies have reported increased rates of disc degeneration in the levels adjacent to spinal fusion. The load of the adjacent levels can be easily assumed to increase, and subsequently, these levels follow the same path. The motion of the adjacent units and their contribution to the overall motion also increases in the unstable phase. Eventually, each cervical unit tends to ankylose with severe degeneration. The mobility of the C4–C5 and C5–C6 U significantly decreases at this stage when compared with the other cervical units and their contribution to the overall motion of the cervical spine significantly decreases.

However, we retrospectively investigated these analyses and did not consider this hypothesis in a persistent model. Thus, using the present investigation as a preliminary study, further prospective research should be undertaken to elucidate the details of the natural history of cervical spine motion following degeneration. Additionally, we believe that kinetic MRI can serve as one of the most useful tools in the prospective research following the investigation of subjects with cervical degeneration.

In conclusion, this study demonstrated the changes in translational motion and angular variation that occur with progressive degeneration. The motion changes with disc degeneration from the normal state to an unstable phase and subsequently to an ankylosed stage with increased stability. We evaluated the contribution of different levels to the changes in overall motion that occurs with degeneration. However, the natural history in cervical spine motion following degeneration has not been elucidated. Further studies are required to provide appropriate treatment methods for the cervical degenerative disease.

### Table 6. Contribution of Each Level to Total Angular Mobility (Percentage)

<table>
<thead>
<tr>
<th>Grade</th>
<th>Level C2–C3</th>
<th>Level C3–C4</th>
<th>Level C4–C5</th>
<th>Level C5–C6</th>
<th>Level C6–C7</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>12.35 ± 8.31</td>
<td>11.95 ± 12.65</td>
<td>21.97 ± 5.78</td>
<td>20.61 ± 12.84</td>
<td>15.40 ± 9.86</td>
</tr>
<tr>
<td>III</td>
<td>12.32 ± 7.97</td>
<td>18.92 ± 9.88*</td>
<td>18.57 ± 12.00</td>
<td>18.60 ± 10.01</td>
<td>15.85 ± 9.88</td>
</tr>
<tr>
<td>V</td>
<td>NA</td>
<td>NA</td>
<td>14.54 ± 9.10*</td>
<td>12.42 ± 9.04*</td>
<td>16.53 ± 11.47</td>
</tr>
</tbody>
</table>

Values are expressed mean ± SD.

*P < 0.05 compared to Grade I at same level.

### Key Points

- Kinetic MRI shows the changes of translational motion and angular variation following degeneration.
- The degenerative process affects the mobility of the functional spinal unit, which moves from a normal disc to a more unstable phase with increased mobility with further degeneration. However, as the degeneration enters the later phases and becomes severe, the motion stabilizes to a more ankylosed stage.
The segments of C4–C5 and C5–C6 U contributed the majority of the total angular mobility of the cervical spine on normal grade. But the roles of C4–C5 and C5–C6 U for total angular mobility significant decreased on severe degeneration.

References

The Effects of the Degenerative Changes in the Functional Spinal Unit on the Kinematics of the Cervical Spine

Yuichiro Morishita, MD, PhD,* Shinichi Hida, MD, PhD,† Masashi Miyazaki, MD,* Soon-Woo Hong, MD, PhD,* Jun Zou, MD,* Feng Wei, MD, PhD,* Masatoshi Naito, MD, PhD,† and Jeffrey C. Wang, MD*

Study Design. The sagittal kinematics of the cervical spine was evaluated using kinematic magnetic resonance imaging (kMRI).

Objective. To investigate the effect of degenerative changes in the functional spinal unit on cervical kinematics by using kMRI.

Summary of Background Data. Few studies have, thus far, by using MR images, described the contribution of degenerative changes in the functional spinal unit to cervical kinematics; however, the exact cervical kinematics remains uncertain.

Methods. A total of 289 consecutive symptomatic patients underwent dynamic cervical MRI in flexion, neutral, and extension postures. All digital measurements and calculations of the variations in segmental angular motion were automatically performed by an MR analyzer using true MR images with 77 predetermined points marked on each image. Each segment was assessed based on the extent of intervertebral disc degeneration (Grades 1–3) and cervical cord compression (groups A–C) observed on T2-weighted MR images.

Results. The segmental mobility of the segments with severe cord compression and moderate disc degeneration tended to be lower than that of the segments with severe cord compression and severe disc degeneration, and a significant difference was observed in the segmental mobility of the C5–C6 segment. Moreover, in all segments with moderate disc degeneration, the segmental mobility was significantly reduced in the presence of severe cord compression, as compared with no compression. However, in segments with severe disc degeneration, no significant differences were observed between the segmental mobility of the cord compression groups.

Conclusion. Our results suggest that cervical cord compression may cause deterioration of cervical cord function and kinematic changes in the cervical spine. We hypothesize that the spinal cord may potentially protect its functions from dynamic mechanical cord compression by restricting segmental motion, and these mechanisms may be closely related to the subject of this manuscript.

Key words: kinematic MRI, functional spinal unit, cervical cord compression, intervertebral disc degeneration, cervical kinematics. Spine 2008;33:E178–E182

The cervical spine is the most mobile region of the spine, affording a wide range of motion. The human spine is subjected to large compressive preloads during activities of daily living. The cervical preload approaches 3 times the weight of the head because of the muscle coactivation forces involved in balancing the head in the neutral posture. The compressive preload on the cervical spine increases during flexion, extension, and other activities of daily living, and is estimated to reach 1200 N in activities involving maximal isometric muscle effort.1

A spinal motion segment is the smallest functional unit of the osteoligamentous spine and exhibits the generic characteristics of the spine. A functional spinal unit (FSU) consists of 2 adjacent vertebrae, the intervertebral disc, and the spinal ligaments (with the exception of the C1–C2 segment). Degenerative changes in the structures of the FSU may ultimately affect the mechanical properties of spinal motion and cause instability and clinical symptoms. Degenerative processes are most prevalent in the C5–C6 segment, followed by C6–C7 and C4–C5.2–4

It may be important to consider the contribution of various factors, such as patient age, gender, neck geometry, degree of degeneration in the cervical spine, history of trauma, and other factors to cervical kinematics. The functional examination of the human spine during flexion–extension along with the measurement of segmental motion in the sagittal plane is a valuable method for analyzing the biomechanics of the human spine. A number of experimental studies on the radiographic and cineradiographic examination of kinematics of the human spine have recently been reported.5–11 However, to the best of our knowledge, few reports have thus far, have based results on kinetic magnetic resonance imaging (kMRI), or described the relationship of the degenerative changes and cord compression in the FSU to the kinematics of the cervical spine. This study examined cervical degenerative changes, such as disc bulging, osteophyte formation, and hypertrophy of the ligamentum flavum, with particular focus on cervical cord compression to evaluate the contributions of these factors to the sagittal plane kinematics of the cervical spine. This study used kMRI to study these variables and their relationship between the effect of degenerative changes in the FSU on cervical kinematics.
Materials and Methods

From February 2006 to May 2007, 289 symptomatic patients (125 men and 164 women) with an average age of 44.0 years (range, 16–93 years) were examined. The subjects consisted of consecutive patients who had neck pain with or without neurogenic symptoms induced by cervical spondylosis. All patients underwent cervical kMRI; scanning was performed with dynamic motion of the cervical spine, including flexion (−40°), neutral (0°), and extension (20°) postures. None of the subjects had previously undergone spinal surgery. The Institutional Review Board approved the study.

MRI Technique

MRI of the cervical spine was performed on a 0.6 Tesla MRI scanner (Upright Multi-Position; Fonar Corp., NY, NY) using a flexible surface coil. The imaging protocol included sagittal T1-weighted spin-echo sequences [repetition time (TR)/echo time (TE), 671/17 milliseconds; slice thickness, 3.0 mm; field of view, 24 cm; matrix, 256 × 200; and number of excitations (NEX), 2] and T2-weighted fast spin-echo sequences (TR/TE, 3432/160 milliseconds; slice thickness, 3.0 mm; field of view, 24 cm; and NEX, 2). All sequences were acquired without fat saturation.

MRI Analysis

The data obtained from the MR images were recorded on a computer for subsequent measurements, and all calculations were automatically performed on true MR images by an MR analyzer. Sagittal MR images were analyzed in 3 positions—flexion, neutral, and extension. For digitization, 77 points were marked on the images by spine surgeons (C3–T1: 6 points on each vertebral body, 2 on each pedicle, and 2 on the spinal canal diameter at each intervertebral disc level; C2: 1 point on the tip of the odontoid process and 6 on the vertebral body; C1: 4 points on the anterior, superior, and inferior surfaces of the anterior tubercle and the lower end of the spinous process; and the occiput (Oc): 2 points on the anterior and posterior baselines) (Figure 1).

The sagittal angular motion was measured for each segment at 5 cervical intervertebral disc levels—C2/3, C3/4, C4/5, C5/6, and C6/7. We defined the total sagittal motion of the cervical spine as the absolute total of the individual sagittal angular motions (C2/3 + C3/4 + C4/5 + C5/6 + C6/7) in degrees, and the contribution of each segment to the total angular mobility of the cervical spine between flexion and extension as percent segmental mobility = (sagittal angular motion of each segment in degrees)/total sagittal angular motion in degrees) × 100.

Assessment of Degenerative Changes in the Cervical Spine

A comprehensive grading system for cervical disc degeneration was obtained by modification of the previously reported system of classification of cervical intervertebral disc degeneration that was based on the degenerative changes in the FSU.12–16 Accordingly, neutral-position T2-weighted sagittal images of 1445 cervical intervertebral discs of 289 subjects were classified into 3 grades (Table 1) by the primary author and were judged eligible for inclusion into the study.

Assessment of Cervical Cord Compression

Table 1. The Grading System for Cervical Intervertebral Disc Degeneration

<table>
<thead>
<tr>
<th>Grade</th>
<th>Nucleus Signal Intensity</th>
<th>Disc Height</th>
<th>Structure of FSU</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Hyperintense</td>
<td>Normal</td>
<td>Without disc herniation</td>
</tr>
<tr>
<td>2</td>
<td>Intermediate/hypointense</td>
<td>Normal/decreased</td>
<td>With/without disc herniation</td>
</tr>
<tr>
<td>3</td>
<td>Hypointense</td>
<td>Decreased/collapsed</td>
<td>With disc herniation/osteophyte</td>
</tr>
</tbody>
</table>

was rated on a 3-point scale (range, 0–2) in which 0 indicated no cervical cord compression, 1 indicated anterior or posterior cervical cord compression not affecting cord alignment, and 2 indicated anterior or posterior cervical cord compression affecting cord alignment. Based on this scale, we classified individual segments into 3 groups: group A, a total of 0 points for each segment; group B, a total of 1 point for each segment; and group C, a total of more than 2 points for each segment. We excluded the C2/3 segment because a few subjects showed cervical cord compression at this level.

Statistical Analysis

Mann-Whitney U test was used for statistical analyses. A P value of less than 0.05 was considered statistically significant.
## Results

The percent segmental mobility for each grade of cervical intervertebral disc degenerative change in each of the segmental cervical cord compression groups is shown in Table 2.

In group A, the grade of degenerative changes in the intervertebral discs increased significantly with an increase in age. With regard to the C3–C4 segment, the average percent segmental mobility at C3–C4 gradually increased with an increase in the grade of degenerative changes in the intervertebral discs, and when compared with Grade 1 changes, significant differences in percent segmental mobility were observed with both Grade 2 and Grade 3 changes (P < 0.01 for Grade 2 and P < 0.05 for Grade 3). With respect to the C4–C5 segment, there was a significant increase in the average percent segmental mobility at C4–C5 with Grade 2 changes (P < 0.05); moreover, percent segmental mobility with Grade 3 changes was almost identical to that with Grade 2 changes. Regarding the C5–C6 segment, the average percent segmental mobility at C5–C6 showed a slight increase with Grade 2 changes and a slight decrease with Grade 3 changes, when compared with Grade 1 changes; however, no significant differences were observed between these values. Regarding the C6/7 segment, the average percent segmental mobility at C6–C7 was almost identical in all groups. In group B, all segments with Grade 3 changes except C4–C5 tended to be associated with a higher age when compared with the segments with Grade 2 changes, and significant differences with respect to age were observed between Grade 2 and Grade 3 changes in both C3–C4 and C4–C5 segments. With respect to all segments, the average percent segmental mobility tended to be lower with Grade 2 changes than with Grade 3 changes; further, a significant difference in the average percent segmental mobility was observed only in C5–C6 (P < 0.05).

Regarding Grade 2 changes, in all segments, there were no significant differences between the groups with respect to age. The average percent segmental mobilities of all segments in group B were almost identical to those in group A; however, the average percent segmental mobilities of all segments in group C were significantly lower than those in group A (P < 0.05 for C3–C4, P < 0.01 for C4–C5, P < 0.001 for C5–C6, and P < 0.05 for C6–C7). In group C, Grade 3 changes in only C4–C5 were associated with a significantly higher age. No significant differences in the average percent segmental mobilities were observed between the groups with respect to all segments with Grade 3 changes.

## Discussion

Many studies have described normal or abnormal cervical kinematics by using different imaging techniques and measurements. Overall, segmental motion of the cervical spine is the least at the C2–C3 level and greatest at the C4–C5 and C5–C6 levels.

The earliest lesions related to degenerative processes of the human spine are thought to occur in the intervertebral disc. Intervertebral disc degeneration typically begins to appear in the second decade of life in men and in the third decade in women, and more than 50% of the...
middle-aged population shows some evidence of cervical spondylosis. 

Because of altered mechanical function of the disc, degenerative changes also begin to occur posteriorly in the facet joints. This degenerative process of the FSU can lead to localized segmental instability or stiffening within different levels of an individual spine. 

Analysis of segmental motion of the cervical spine may help in detecting degeneration or damage within the spine.

Regarding the effect of dynamic motion on the cervical spinal cord, the cervical cord shortens and its cross-sectional area increases during extension of the cervical spine; however, during flexion, it stretches, leading to increased axial tension. These mechanical stresses on the cervical cord as well as static factors, such as disc herniation, osteophyte formation, and hypertrophy of the ligamentum flavum that result from degenerative changes in the FSU, contribute to the pathogenesis of cervical spondylitic myelopathy.

Chen et al. reported that increased segmental angular motion may reduce the sagittal diameter of the spinal canal and lead to spinal canal stenosis associated with disc bulging and hypertrophy of the ligamentum flavum. In contrast, Miha et al. focused on canal stenosis at the C3–C4 level and reported that in elderly patients with cervical spondylitic myelopathy due to canal stenosis at the C3–C4 level, the C3–C4 segmental angular motion was significantly greater than that in younger subjects or in the elderly healthy population. They hypothesized that an age-related reduction in the mobility of the lower cervical segments may promote mechanical stresses on the upper cervical segments, leading to canal stenosis at the C3–C4 level. However, they discussed only the process of cervical spinal canal stenosis in relation with the degenerative changes in the cervical spine. We presumed that the next change in cervical kinematics might occur after the formation of cervical spinal canal stenosis with degenerative changes in the cervical spine.

It is generally accepted that there are 3 separate stages of clinical manifestations of degenerative changes in the intervertebral discs; these include temporary dysfunction, an unstable phase, and stabilization with progression of the degenerative changes. 

In our study, in all segments within all the compression groups, the degenerative changes in the intervertebral discs tended to progress with age. However, in C3–C4 and C4–C5 with no cervical cord compression, the contribution of each segment to total cervical mobility increased with progression of the degenerative changes in the intervertebral discs. By using kinematic MR images, a total of 1445 functional spinal units of 289 symptomatic subjects were examined for intervertebral disc degeneration and cervical cord compression.

On the other hand, in segments with cervical cord compression, particularly severe cord compression, segmental mobility tended to show lower values in segments with moderate intervertebral disc degeneration than in those with severe degeneration; moreover, a significant difference was observed in the segmental mobility at C5/6. Moreover, in all segments with moderate disc degeneration, there were no significant differences between the compression groups with respect to age, and the segmental mobility was significantly reduced in the segments with severe cord compression when compared with those with no cord compression. These results suggest that cervical cord compression greatly affects the sagittal segmental motion of the cervical spine only if there is sufficient intervertebral disc height and flexibility. We hypothesize that the spinal cord may shift horizontally to prevent lesions that develop due to cord compression. However, in severe cord compression that affects spinal cord alignment and causes cord impingement, the spinal cord cannot shift away and escape compression and may be affected by restriction of segmental motion. Cervical cord compression may result in not only deterioration of the cervical cord function but also kinematic changes in the cervical spine. The spinal cord may protect its function from dynamic mechanical cord compression by restricting segmental motion.

However, in segments with severe intervertebral disc degeneration and decreased height or collapse of the intervertebral disc, no significant differences were observed between the segmental mobilities of the cord compression groups. This result suggests that when the intervertebral discs are stiffened due to severe degenerative changes such as disc height loss or osteophyte formation, sagittal segmental motion of the cervical spine is only mildly affected by cervical cord compression. Moreover, we hypothesized that the mechanisms for protecting the spinal cord may be closely related to the intervertebral discs. Deterioration in intervertebral disc function may lead to deterioration in the mechanisms for protecting the spinal cord.

However, some issues remain unanswered even in the current study. We did not discuss the relationship between cervical kinematics and other factors such as gender, age, cervical alignment, and clinical manifestations. Therefore, using the current investigation as a pilot study, further research using larger patient populations may help to resolve several unclear results obtained from this study; moreover, the details of the relationship between cervical kinematics and the degenerative changes in the FSU can be clarified further.

### Key Points
- By using kinematic MR images, a total of 1445 functional spinal units of 289 symptomatic subjects were examined for intervertebral disc degeneration and cervical cord compression.
In segments with severe cord compression and moderate disc degeneration, the segmental mobility tended to have lower values than in those with severe cord compression and severe disc degeneration.

In all segments with moderate disc degeneration, the segmental mobility was significantly reduced in the presence of severe cord compression when compared with no compression; however, in segments with severe disc degeneration, no significant differences were observed between the cord compression groups with regard to segmental mobility.

Our results suggest that cervical cord compression may result in not only deterioration of cervical cord function but also kinematic changes in the cervical spine.

We hypothesize that the spinal cord may potentially protect its functions from dynamic mechanical cord compression by restricting segmental motion, and these mechanisms may be closely related to the intervertebral discs.

References