Ex Vivo Biomechanical Testing to Examine the Etiology of Low Back Pain as a Result of Whole Body Vibration

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### Abstract

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An association between occupational whole body vibration (WBV) exposure and the development of lower back pain (LBP) has been established through numerous epidemiological studies. However, the etiological and biomechanical mechanisms of how WBV contributes to LBP and injury are not well understood. Using field-measured occupational WBV exposures in a controlled laboratory setting, this study attempted to expose the human lumbar spine to occupational WBV exposures to better understand the mechanisms of WBV-related low back injury. The occupational exposures selected for this study were those encountered by bus drivers, who are known to have a high prevalence of LBP. Continuous and impulsive WBV accelerations collected at the floor of a bus traversing a standardized route in the Seattle metro area were played into a six-degree of freedom hydraulic shaker platform. Twelve professional drivers sat on an air suspension seat mounted to a six-degree of freedom hydraulic shaker platform. The drivers' response to the continuous and impulsive exposures at the seat and sternum were

recorded. The difference between seat- and sternum-measured accelerations were used to calculate the displacement of the spine during the vibrations encountered during continuous and impulsive exposures. Based on these two exposure patterns for the bus drivers, six human lumbar functional spine units (FSUs) were placed in a servohydraulic testing system and continuously exposed to one of the two vibration signatures. A unique staining technique was used to visually differentiate preexisting intervertebral disc damage and damage brought on by cycling the FSU. No discernible damage was found for discs exposed to the continuous WBV while exposing FSUs to the impulsive WBV resulted in endplate fractures and damage to the annulus fibrosis of the intervertebral disc. These findings indicate that, for equal drive times, a more impulsive and intermittent loading signature is more likely to cause injury to components of the low back than a relatively less impulsive and more continuous signature.

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#### **CHAPTER 1: Background and Introduction**

#### 1.1 Relationship between Whole Body Vibrations and Lower Back Disorders

Excluding head injuries, back injuries are the most prevalent injury experienced by adults in the United States (US) <sup>1</sup>. Back injuries and lower back disorders (LBDs) are also the most prevalent and costly non-lethal medical condition affecting the US workforce. The lifetime prevalence rate of LBDs in US adults is nearly 80% and total annual costs of treatment exceed 100 billion dollars <sup>2, 3</sup>. Additionally, back injuries account for nearly one-third of all work-related injuries in the United States and one-fifth of all US workers' compensation claims, thus carrying significant financial implications <sup>3</sup>.

Workplace LBD etiology is believed to be multi-factorial in its origin and related to physical and psychosocial occupational exposures <sup>4-6</sup>. Amongst sedentary workers and in occupational settings requiring prolonged periods of sitting, both physiologic deconditioning and continuous and prolonged static musculoskeletal loading are believed to play a role in the onset and development of LBDs <sup>4</sup>. Additionally, poor workstation ergonomics and awkward sitting postures have been found to increase the likelihood of LBD onset amongst occupational groups with prolonged sitting requirements <sup>6,7</sup>.

Machine and vehicle operators have higher rates of LBD prevalence compared to similar groups of office workers, despite the sedentary nature of both types of occupations <sup>8-10</sup>. It is believed that exposure to whole body vibrations (WBV) such as those experienced during vehicle operation are a major factor in this increased prevalence and a leading risk factor in the onset and development of LBDs in the workplace <sup>3, 11-14</sup>. WBV in vehicles is primarily transferred to the operator via contact surfaces supporting the body, such as the buttocks of the seated individual. In regards to LBDs, WBV can be traumatic for individuals as vertebral

structures (muscles, ligaments, vertebral bodies, and intervertebral discs) act as springs and dampers to dissipate vibration energy <sup>11, 15</sup>. This vibration energy dissipation is particularly pronounced in the lumbar region of the spine, where both elevated and repetitive spinal loads have been associated with vertebral disc thinning, bulging, and herniation <sup>12, 16, 17</sup>. Additionally, cyclical torsion and bending has been found to lead to disc degeneration and micro-fractures in the vertebral bodies of the lower back <sup>18</sup>. Because of this, degenerative lumbar discs are 150% more likely in vibrating vehicle operators exposed to high levels of WBV than those that are less exposed <sup>3</sup>. It is believed that 4-7% of the US workforce is exposed to potentially harmful levels of WBV, among which 41-83% of vibrating vehicle operators report lower back pain as a result of their occupation <sup>12, 15</sup>. High levels of both WBV and LBDs have been reported in operators of cars, buses, trucks, trains, forklifts, all-terrain vehicles, mine cars, tractors and other agricultural equipment <sup>13, 19-29</sup>. Thus, vehicle operators are believed to not only be at risk for development of LBDs due to the sedentary nature of their work, but also because of an increased exposure to WBV.

## 1.2 Brief Overview of Spine Anatomy

As can be assumed from the word "back", most LBDs are in some way related to the vertebral (spinal) column and/or the surrounding musculature thereof. The spinal column consists of 33 vertebra and 26 bones grouped into five sections/regions <sup>30</sup>. Figure 1.1 shows a diagram of these structures and their groupings. From the superior (top) of the vertebral column, there are 7 cervical vertebra, 12 thoracic vertebra, and 5 lumbar vertebra <sup>30</sup>. Each of these 24 vertebrae are composed of individual bones stacked one atop another along the cranial-caudal axis with intervertebral discs between adjacent vertebrae. Twenty-three discs are contained in the

human spine – 22 between each adjacent vertebra and 1 immediately inferior to the lumbar spine at the lumbo-sacral junction. The vertebra are situated to create four distinct, curved regions with the sacral and thoracic curves present at birth and the lumbar and cervical curves developed thereafter <sup>30</sup>. When classifying individual vertebrae, a single letter is used along with a number representing the location of the vertebra of interest. These letters correspond to the first letter of the region from which the vertebra originates while the numbers correspond to the individual vertebra's location from the superior for the particular region of interest. As examples, the superior-most cervical vertebra would be C1, the 5<sup>th</sup>-most superior thoracic vertebra from would be labeled T5, and the 2<sup>nd</sup>-most superior lumbar vertebra would be L2. When fused, the sacrum is collectively referred to as S1.



Figure 1.1 Overview of vertebral column structures.

The vertebrae are the primary supporting structures of the vertebral column while the

intervertebral discs provide flexibility and shock absorption. Individual vertebrae are composed of a hard cortical bone cortex and a cancellous (trabecular) bone interior. Stabilizing structures are found protruding from the posterior regions of vertebrae. Meanwhile, intervertebral discs are connected with vertebrae (above and below) at the vertebral "endplates." The discs are composed of an outer annulus fibrosis surrounding an inner nucleus pulposus. The layout of intervertebral discs is outlined in Figure 1.2. The annulus fibrosis (annulus) is composed of collagen fibers and keeps the nucleus pulpsous (nucleus) contained. The nucleus is composed of a protein-based hydrogel functioning as a shock absorber.



Figure 1.2 Anatomy of intervertebral discs.

Injuries to the spine and back can take many forms. The nucleus is free to move within the area confined by the inner annulus fibers – as loads are applied to the vertebral bodies adjacent to the disc, a pressure gradient is created within the inner annulus which forces the nucleus away from the location of the load. Loading vertebral structures along the cranial-caudal axis also forces the annulus fibers to expand along the transverse plane and into the space between

vertebral bodies. Coupled with the movement of the nucleus towards this same region, this can lead to disc bulging, herniation, or impinging of the nerves running along the posterior of the vertebral bodies. In cases of extreme loading, endplate fractures can also occur. Laboratory tests have shown that disc bulging, rupturing of the annulus, and endplate fractures can occur during tasks which mimic occupational lifting situations <sup>31</sup>. However, it should be noted that in general, back pain is often grouped nonspecifically when classifying workplace injury, thereby making it difficult to determine the nature of occupational back injuries as brought on by WBV exposure or otherwise <sup>32</sup>.

# 1.3 Load-tolerance and Variability Within the Spine

Biomechanically, LBDs are associated with the magnitude and frequency of lumbar spine loading. Increased loading results in a decrease in the spine's ability to tolerate applied loads. The related load-tolerance relationship is graphically shown in Figure 1.3<sup>-1</sup>. Central to this relationship is the hypothesis that if the load experienced by a region of the vertebral column exceeds a physiologically-specific tolerance, the vertebral tissue of that region may be subject to deterioration and/or damage. Figure 1.3A and 1.3B depict this relationship for a spinal load significantly less than the tolerance level (as indicated by a "safety margin") and for a load magnitude greater than the tolerance level, respectively. Figure 1.3C depicts this relationship for cumulative trauma disorders, where spinal load tolerance decreases with non-uniform, repetitive loading over time <sup>1, 33</sup>. This gradual decrease of spinal load tolerance over time is representative of the commonly held view that the development of WBV-induced LBDs are believed to progress in a manner similar to cumulative trauma disorders – spinal tolerance decreases over time and realization of injury can often be highlighted by a single event.



Figure 1.3 Load-tolerance curves (1). 1.3A: Tolerance greater than load. 1.3B: Load greater than tolerance. 1.3C: Cumulative trauma from decreased tolerance over time. Reproduced from source.

Even in light of this load tolerance-relationship, the etiological relationship between WBV and LBDs is still relatively unknown<sup>14</sup>. This is largely due to both the complexity of the biomechanically-relevant anatomy of the spine and the complexity of nociception and pain response registered and regulated by the nervous system. In general, back pain has both neurophysiological and neuroanatomical origins but the complexity and anatomic variation within the spine make determining the underlying causes of low back pain difficult <sup>1, 34, 35</sup>. In the spine, both the average height and average width of lumbar vertebrae are two to three times those of thoracic and cervical vertebrae, respectively <sup>36</sup>. Also, the facet joints in the lumbar region of the spine differ from other regions of the spine (particularly in relation to facet joint angle with respect to midline), and within the inner/outer regions of an individual vertebra <sup>1, 36, 37</sup>. Due to slight anatomical variations in the intervertebral discs and the increasing effects of body mass as one goes down the spinal column, studies have indicated that WBV does not uniformly affect all regions and structures of the spine <sup>36</sup>. Eccentric loading is the primary cause of disc herniation and bulging, giving basis to the idea that facet joints (located on the posterior region of vertebrae) and the posterior region of lumbar intervertebral discs, which are exposed to the highest magnitudes of vibration, are at greatest risk for WBV-induced injury <sup>36</sup>. As such, only a strong epidemiological association between WBV exposure and LBDs exists while etiological evidence regarding the nature, progression, and location of injury remains lacking.

#### 1.4 Previous Work

Previous WBV studies have focused on four broad areas: conducting epidemiological surveys to establish an association between exposure and disease, collecting field data with respect to established standards in hopes of developing engineering controls, modeling seat suspension performance using laboratory shaker platforms, and performing laboratory simulations using finite element (FE) or cadaveric models. Data for WBV studies is almost always collected and analyzed in accordance with International Organization for Standardization (ISO) guidelines using accelerometers and weighted acceleration-based parameters <sup>38, 39</sup>. These parameters include, among others, a time-weighted average acceleration (A<sub>w</sub>) used to measure the continuous component of the vibration exposure; a fourth-order time-weighted acceleration (VDV) to better account for the impulsive content of the vibration exposures; and a high fidelity, continuous acceleration measure (S<sub>ed</sub>) to better evaluate and model the impulsive, compressive stresses in the lumbar region of the spine <sup>38, 39</sup>. It should be noted, however, that some of these parameters and the coefficients used in their calculations may be outdated, may not appropriately identify harmful levels of WBV, and the calculations of the various parameters may be subject to errors in interpretation <sup>40-43</sup>. Additionally, occupational WBV is poorly understood and regulated in work settings as studies have recorded occupational WBV exposures significantly greater than ISO and European Union Directive risk guidelines <sup>12, 19, 25, 28, 44</sup>.

Approaches to characterize the work-relatedness of WBV-induced injuries have focused on establishing a relationship between WBV and LBDs. This has primarily been done via crosssectional, cohort, case-control, and community-based epidemiological studies of WBV-exposed groups <sup>12</sup>. Crane, bus, tractor, and fork-lift operators have been frequently studied for both vibration exposure and LBDs with control groups usually consisting of office workers <sup>12</sup>. In general, such studies have shown that not only does daily WBV exposure in many industrial vehicles exceed industry-standard exposure limits, but WBV is also a risk factor for LBD based on the higher prevalence rates in the studied populations <sup>12</sup>. Even though the association between WBV and LBD is believed to be based on some combination of mechanical overloading and muscular fatigue, the exact biological etiology of WBV-induced LBD remains relatively unknown <sup>14</sup>. Additionally, many variables can affect the level of WBV exposure including seat design, hand position, driving posture, and vehicle design <sup>11, 22, 29, 42, 45, 46</sup>.

Computational simulations and analyses have been conducted to investigate potential mechanisms of lower back injury relating to WBV. In particular, FE models have provided researchers with a tool by which to accurately assess the biomechanics of the lumbar spine under a range of conditions, albeit under the limitations posed by replacing cadaveric experimentation with computational simulations <sup>47-50</sup>. One such FE analysis found that both static and dynamic compressive and sheer forces acting on lumbar bodies during whole body vibrations were highly dependent on upper body mass and posture <sup>51</sup>. A transfer function analysis of the sheer and compressive forces, meanwhile, found these forces to be highest when the frequency of vibration is under 10-12 Hz in the z-direction (caudal-cranial direction when subject is seated upright) <sup>51</sup>. Another FE analysis found the resonant frequency of the lumbar spine under vibration to vary greatly based on upper body mass but to be about 10 Hz in the z-direction <sup>33</sup>. However, the health of intervertebral bodies, posture, and body mass was found by another FE analysis to have a large influence on the resonant frequency of lumbar vertebral bodies <sup>52</sup>. Both a larger upper

body mass and a higher deterioration/removal of vertebral bodies were found to decrease the resonant frequency of lumbar vertebral regions in a z-axis vibration <sup>50, 52</sup>. Static, peak compressive forces acting on the lumbar spine, meanwhile, have been found by FE analyses to range between about 350 N-700 N when the subject is seated in an ergonomically suitable driving posture with a 40 kg upper body mass; when the subject is bent slightly forward, these compressive forces have been found to increase to between about 650 N-1200 N <sup>43, 51</sup>. Thus, the number of variables that have been found to influence WBV resonant frequencies and the forces acting on vertebral bodies vary largely across occupational settings and are both vehicle- and operator-dependent.

FE analyses have also been useful in determining which regions of the lumbar spine are most susceptible to deterioration when exposed to WBV. Under cyclic, sinusoidal loading, it was found that posterior regions of lumbar vertebral bodies are subjected to greater forces under WBV than anterior regions <sup>52</sup>. Interestingly, this is in opposition to the widely-held view regarding natural lumbar degeneration, which is believed to proceed from the anterior region towards the posterior <sup>53</sup>. Additionally, FE analyses have indicated that facet joints on lumbar vertebral bodies may be of increased risk of deterioration when dynamically exposed to vertical WBV compared to static loading <sup>52</sup>. Facet contact forces were also found to increase with increasing frequency as well as with intervertebral disc degeneration and/or removal <sup>54</sup>.

Cadaveric experiments have also been conducted to determine the effects of WBV on vertebral segments. One such experiment determined individual lumbar segments to have resonant frequencies near 5 Hz <sup>55</sup>. Intradiscal pressure was also found to be greatly dependent on posture, increasing when leaning forward <sup>56</sup>. Seat to vertebrae vibration and pressure transmission in these cadaveric experiments was found be highest at about 5-7 Hz <sup>56, 57</sup>. In

analyzing these cadaveric results, it must be noted that many biological functions, not the least of which are muscle contraction and thermostasis, which could alter vibration characteristics are not present <sup>56, 57</sup>.

### 1.5 Problem Statement and Proposed Solution

As discussed previously, LBDs are amongst the most common occupational health hazards. A leading risk factor for the onset and development of LBDs in the workplace is WBV. Though an exposure-response relationship between WBV and LBDs has been well established, the exact causal pathway for onset of LBDs from WBV is not clearly understood. This is both due to a wide array of ergonomic and occupational variables associated with vehicle-related WBV and due to the anatomic complexity of the spine.

Based on previous work, the causal relationship between WBV and LBDs has been well established; therefore, attempting to determine the etiological implications thereof is one logical, next step in better understanding occupational WBV as a whole. Collecting occupational WBV exposures from real-world settings, using this data to apply realistic, occupational vibration signatures to human spinal segments, and attempting to analyze and characterize the impact of the biomechanical loading on the tissues could be of great utility in understanding the relationship between WBV and LBDs. The experiment upon which this thesis is based used field-collected WBV signatures to impart occupational vibration exposures on cadaveric tissues. Occupational vibration exposures were collected in King County Metro buses, processed, and played into a vibration shaker platform upon which subjects were seated. Movement of the subjects' sternum relative to the seat was then calculated for these subjects and these calculated displacements were then used to guide ex vivo loading experiments of intervertebral discs. By relating WBV to true-to-life exposures, hopes are that these experiments will be used to better understand the physiological and biomechanical implications of occupational WBV on LBDs, particularly with respect to different types of vibration exposures.

This may be one of the first experiments to use occupationally collected WBV exposures to impart biomechanical loads on human cadaveric functional spinal units in a controlled laboratory setting. As such, no known protocols and procedures were in place prior to the start of this project. Additionally, previous studies outlining disc injury as a result of repeated cycling relied on radiography to determine the extent of disc injury or migration of nuclear material into the annulus of the intervertebral disc  $^{58-60}$ . This project sought to develop an alternative and more cost-effective means to outline progression of disc deterioration and damage. As stated previously, the nature of occupational disc injuries has not been well documented and the progression thereof is not particularly well understood. Specifically, it is not known if progression of injury as a result of WBV exposure extends from the exterior of the intervertebral disc towards the interior or progresses from the interior of the disc towards the exterior. Prior studies examining the loading responses of vertebral segments have identified an outwardlyprogressing disc pathology, however, the discs' response to WBV signatures is of yet unknown <sup>58, 59</sup> For this project, a testing protocol was developed based on the assumption that injury progresses away from the most-pressurized region of the disc, i.e. away from the location of nucleus pulposus. As such, an attempt was made to verify that this assumption is in fact valid without necessarily bringing the validity of the view that disc injury progresses inwardly into question.

Based on the above, the objectives of this experiment were as follow:

1. Develop a protocol by which occupational WBV exposures can be translated into a

laboratory setting for testing on human cadaveric specimens.

- Develop a non-radiographic protocol by which injuries induced during laboratory testing can be traced.
- 3. Determine which of two exposures one with a higher amplitude of vibration and one with a more continuous cycle of vibrations is comparatively more injurious to the lower back.
- 4. Determine what types of injuries and/or damage to the vertebral column can be seen as a result of the different spinal loading protocols.
- 5. Determine the progression of disc related deterioration, if any is induced during cycling.
- 1.6 Thesis Overview

This thesis consists of 6 chapters and is organized as follows. Chapter 2 outlines the process by which occupational exposure data was collected and analyzed. Chapter 3 details the methods and results of preliminary ex vivo experimentation and revisions to the testing protocol. Chapter 4 discusses results from ex vivo experimentation after finalizing a testing protocol from the preliminary tests. Chapter 5 discusses major findings from Chapter 4, summarizes all pervious chapters, and details limitations of the experiment as well as improvements to be considered for a second-generation of experiments.

#### **CHAPTER 2: Vibration Data Collection and Processing**

# 2.1 Vibration Data Collection and Processing Overview

This chapter discusses the methods and protocols used in obtaining WBV signatures to be used in subsequent laboratory testing on human cadaveric specimens. Occupational WBV data was collected in King County Metro buses at the vehicle seat and floor by Ryan Blood, a Ph.D. student in the University of Washington's Department of Environmental and Occupational Health Sciences (henceforth referred to as "Blood"). Because sternum vibration data was not collected during this initial round of data collection, the data from the buses was to be used in conjunction with a computational mathematical model to predict the response of the drivers' sternums to vibrations. Blood later collected vibration data at drivers' sternums using a hydraulic shaker platform using the bus data as inputs. This data was processed to extract the displacement response of individual intervertebral discs to vibrations and verified using the computational mathematical model. This extracted data was used in the laboratory experiments detailed in later chapters. An overview of the process described in this chapter is shown in Figure 2.1.





# 2.2 Bus and Shaker Data Collection

Occupational WBV data was first collected in the field WBV data and then the WBV data was inputted into a six-degree of freedom hydraulic shaker platform. This input data for the shaker platform as well as the data from the shaker platform were collected by colleagues. What follows is a description of the data collection process as outlined by Blood who helped coordinate both cycles of data collection. Methods from the bus data collection have previously been published <sup>3</sup>.



Figure 2.2 Standardized route used in data collection.

WBV field data was collected on a bus (13.9m long bus; New Flyer; Winnipeg, Manitoba, Canada) as a driver traversed a standardized route spanning the greater Seattle area

with no passengers other than the driver and at most two research staff. The route consisted of a variety of road conditions including city streets, an older freeway segment (Interstate 5; I5), a newer freeway segment (Interstate 90; I90), and a segment with speed bumps. A map of the route is provided in Figure 2.2 and coordinates of the road segments are provided in Table 2.1.

Segment		Latitude	Longitude	Direction of Travel
City	Start	47° 30' 30'' N	122° 17' 28" W	Northwest
(12 km)	End	47° 35' 25" N	122° 19' 46" W	East
New	Start	47° 35' 36" N	122° 19' 19" W	East
(29 km)	End	47° 34' 44" N	122° 8' 47" W	East
Speed	Start	47° 35' 20" N	122° 8' 44" W	West
(1 km)	End	47° 35' 7" N	122° 8' 36" W	West
Old Freeway (10 km)	Start	47° 35' 27" N	122° 19' 18" W	South
	End	47° 30' 35" N	122° 17' 1" W	Southeast

 Table 2.1
 GPS coordinates of road segments in standardized route

Data was collected on the bus using a four-channel data recorder (model DA-20; Rion Co.; Tokyo, Japan) logging both GPS data and accelerometer data at 1 Hz and 1280 Hz, respectively <sup>3</sup>. The GPS recorder (Model DG-100; GlobalSat; Chino, CA) was mounted in the cabin of the vehicle and used to determine vehicle location and velocity. Acceleration data was acquired in the z-axis (perpendicular to the floor of bus) at the vehicle floor using a tri-axial accelerometer (model 356B40; PCB Piezotronics; Depew, NY) magnetically mounted to the floor of the bus under the driver's seat. Collected data was stored on a 2 gigabyte compact flash memory card (Extreme III; San Disk; Milpitas, CA). GPS and acceleration data were

synchronized during post-processing. The accelerometer was calibrated prior to data collection and had a working bandwidth between 0.5 Hz and 1 kHz.

Initially, the experimental plan was to use the vibration data from the bus seat in conjunction with a computational mathematical model to predict the response of the torso and sternum to vibrations. This mathematical model is detailed in section 2.4 of this writing. The mathematical model was to be relied upon because only vibration accelerations at the seat and floor of buses had been collected – the driver was not instrumented with accelerometers or other instrumentation that could measure and detect the spine's response to vibrations. However, after the seat vibration data was collected in the field setting, Blood subsequently collected WBV data when subjects sat in the same seat used in the bus field data collection and were subjected to the field-collected vibrations using a hydraulic shaker platform. In addition to measuring the floor and seat measured accelerations while on the shaker platform, subjects were also outfitted with a tri-axial accelerometer mounted over the sternum. This sternum-collected vibration data was processed for the experimental cycling protocol and the results were compared with the computational mathematical model. The processing of results from this data collection is detailed in section 2.5 and the comparison with the mathematical model is described in detail in section 2.4. A brief description of Blood's shaker collection protocol is provided below.

Blood used the floor accelerations from the bus as inputs into a six-degree of freedom hydraulic shaker platform (model Standard; Moog Inc.; Kirkland, WA). In total, 13 different vibration signatures were played into the shaker platform to simulate road and vehicle conditions. Three of the inputs were from the collected bus vibration data. The 13 input signals are detailed in Table 2.2. 12 male subjects were seated in an industry-standard mechanical seat with an air suspension system (model Q91; USSC Seating; Exton, PA) which was securely fixed

to the top of the platform. The 13 input vibrations were generated while the drivers were seated on top of the platform. Tri-axial accelerometers identical to those used in the bus data collection were mounted to the base of the platform, the seat top, and to the chest of each subject, using a magnet, an ICP seat pad, and an elastic band secured to the subject's chest, respectively. The floor accelerometers were used to validate the fidelity of the shaker platform. The experimental setup is illustrated in Figure 2.3 on the next page. An eight-channel data recorder (model CoCo 80; Crystal Instruments; Santa Clara, CA) was used to log the acceleration at a rate of 1280 Hz. Additional information regarding the subjects is provided in Table 2.3.

Signal	From Bus Vibration Data?	Used for Data Processing?	Signal Description	Duration (Seconds)
1			0.1 Hz, 100 mm amplitude sinusoid	40
2			No signal (break 1)	7
3			3 Hz, 3 mm amplitude sinusoid	40
4			No signal (break 2)	7
5			5.5 Hz, 1.5 mm amplitude sinusoid	40
6			No signal (break 3)	7
7			8 Hz, 0.5 mm amplitude sinusoid	40
8			No signal (break 4)	60
9	~		City streets input	180
10			No signal (break 5)	60
11	~	$\checkmark$	Smooth freeway input	180
12			No signal (break 6)	60
13	~	$\checkmark$	Rough freeway input	180

Table 2.2Input signals for shaker table

### Table 2.3Shaker table subject information

	Age (years)	Height (cm)	Weight (kg)
Mean for all subjects (SD)	36.8 (± 9.4)	181.1 (± 9.1)	111.5 (± 34.8)



Figure 2.3 Shaker table experiment setup. Locations of arrows mark approximate placement of three accelerometers (image courtesy of Ryan Blood).

Of the 13 input signals into the hexapod, only the smooth and rough freeway road segments (inputs 11 and 13 in Table 2.2) were used for subsequent cadaveric experiments outlined in Chapters 3 and 4. As can be seen in Figure 2.2, the "smooth freeway input" consisted of data collected along eastbound Interstate 90 near Seattle, WA and the "rough freeway input" consisted of data collected along southbound Interstate 5 near Seattle, WA. The two segments were traversed with similar average speeds:  $88.7 \pm 4.0$  kmph for the smooth freeway and  $93.3 \pm 4.4$  kmph for the rough freeway, as indicated by the speed output collected by the GPS unit. The chief difference between the two segments was that the smooth freeway segment consisted

mostly of even, unpatched road while the rough freeway segment consisted of a freeway with expansion joints. These expansion joints, distanced about 50 m (about 2 seconds in driving time) apart, imparted impulses to the driver that made the otherwise similar road signatures very different. The two segments are further detailed in Table 2.4. The seat and chest acceleration data from the hexapod subjects were converted to displacements and used to analyze vibration signatures for the smooth and rough freeway inputs.

Hexapod Input Signal	Segment		Latitude	Longitude	Direction of Travel
11	Smooth freeway (12 km)	Start	47° 35' 15" N	122° 13' 55" W	South
		End	47° 34' 40" N	122° 12' 28" W	South
13	Rough freeway (29 km)	Start	47° 34' 30" N	122° 19' 12" W	Southeast
		End	47° 33' 15" N	122° 19' 7" W	Southeast

Table 2.4Segments used in post-shaker table analysis

#### 2.3 Triple Filter, Double Integration Approach

#### 2.3.1 Overview of Approach

Standards for WBV data collection are based on the acceleration encountered and not displacement profiles <sup>38, 39, 61</sup>. For the inputs to be used in the proposed experimental cycling to human cadaveric spinal segments, displacement values were needed as required for inputs to the servohydraulic testing equipment used. Therefore, for the cycling experiments, a method was needed to convert the experimental acceleration data to displacement data. Formulaically, the velocity and displacement of a body can be determined from a single and double integration, respectively, of its acceleration <sup>62</sup>:

$$v(t_f) = v(t_0) + \int_{t_0}^{t_f} a(\tau) d\tau \qquad (eq.1)$$

$$d(t_{f}) = d(t_{0}) + \int_{t_{0}}^{t_{f}} v(\tau) d\tau \qquad (eq.2)$$

Where the variables "a," "v," and "d" symbolize acceleration, velocity, and displacement, respectively while "t" represents values in the time domain and " $\tau$ " represents a dummy integration variable. Apparent from both eq. (1) and eq. (2) is that to accurately calculate velocity and displacement from acceleration, initial values of velocity and displacement are needed <sup>62</sup>. When working with discretely collected acceleration data, however, initial values for velocity and displacement cannot be extracted from acceleration data – the acceleration only provides a measure of change in regards to initial velocity which, in turn, only provides a measure of change in regards to displacement. Additionally, formulas such as those in eq. (1) and eq. (2) are extremely difficult to apply to discretely collected data with no formulaic basis (such as the data collected from the buses and the hydraulic shaker platform). Therefore, the method of calculating displacement from acceleration data discussed henceforth does not require knowledge of the initial state of the accelerating body and does not rely on the application of formulas for the conversion of acceleration data.

To calculate displacement from acceleration data, three stages of high-pass filtering were used in conjunction with two stages of integration. First, the raw acceleration data was filtered using a high-pass filter and the filtered acceleration data was integrated to calculate velocity. This velocity data was then filtered using a high-pass filter and the filtered velocity data was integrated to calculate displacement. Finally, the displacement data was filtered using a high-pass filter to produce the desired result. The high-pass filters were used to account for accelerometer drift, DC biases, and the lack of initial conditions. Accounting for DC bias was of particular integrated as any constant gain from a signal gets compounded during integration, thereby causing miscalculations. A block diagram of the triple-filter-double-integration (TFDI) process is shown in Figure 2.4. All integrals in the triple-filter-double-integration process were calculated using a Riemann sum approximation via the trapezoidal rule across each data point. An 8<sup>th</sup> order dualpass Butterworth high-pass filter (4<sup>th</sup> order each pass) with a cutoff of 0.5 Hz was used in filtering. A high filter order was chosen so filter settling time would be minimal and a Butterworth filter was used to have zero gain or loss of signal in the passbands. A dual-pass filter was used so filtering did not introduce a phase shift during any iteration of filtering. High-pass filtering allowed for acceleration data to be integrated numerically without relying on formulas.



Figure 2.4 Overview of triple-filter-double-integration (TFDI) process.

### 2.3.2 Validation of Approach

To determine the effectiveness of the TFDI process in accurately converting acceleration signals to displacement values, a servohydraulic testing system was used. A tri-axial accelerometer identical to that used in prior data collection was magnetically mounted to the actuator of a servohydraulic testing system (model 810; MTS; Eden Prarie, MN). Uniaxial (z-axis) acceleration data was logged using a 8-channel data collection system (model DA-40; Rion Co.; Tokyo, Japan) for about 1 min at 1280 Hz as the MTS's actuator oscillated at two different peak-to-peak amplitudes (0.2 mm and 2 mm) at each of three different frequencies (0.5 Hz, 1.5 Hz, and 3 Hz). This acceleration data was then processed using the TFDI process to extract the

displacement of the actuator. The results from this extraction are shown in Figure 2.5 on the next page (please note that to account for filtering effects and filter roll-offs, the data shown is from the 30 second mark in data collection onwards). For reference, also shown in Figure 2.5 are computer-generated sinusoids of the expected amplitude and frequency.

The calculated displacement data was also processed using a computer software algorithm to detect local minima and maxima of each of the six displacement signals by first numerically finding the derivative of the signal and then finding zero crossings of this derivative. This peak detection algorithm (henceforth referred to throughout this thesis thusly) was used to determine both the peak-to-peak amplitude and the duration between each successive minima and maxima of the calculated displacement data sets. To account for filtering settling effects from roll-offs of the passband, the first 10 seconds of each data set were excluded in calculations and 40 seconds thereafter (50 seconds into the data collection) were used as an end time. The amplitude and time calculations from the peak detection algorithm were then used to determine the accuracy of the TFDI approach, as detailed in Table 2.5.

Run	Expected Frequency	Expected Amplitude	Calculated Frequency	Percent Error Frequency	Calculated Amplitude	Percent Error Amplitude
А	3 Hz	2 mm	3.00 Hz	< 1%	2.00 mm	< 1%
В	1.5 Hz	2 mm	1.50 Hz	< 1%	1.96 mm	2.1%
С	0.5 Hz	2 mm	0.56 Hz	11.9%	0.15 mm	92.5%
D	0.5 Hz	0.2 mm	1.16 Hz	132.8%	0.03 mm	83.8%
Е	1.5 Hz	0.2 mm	1.50 Hz	< 1%	0.19 mm	3.1%
F	3 Hz	0.2 mm	3.00 Hz	< 1%	0.20 mm	< 1%

 Table 2.5
 Results of TFDI validation calculations



Figure 2.5 TFDI calculations and validation. A: 3 Hz, 2 mm amplitude; B: 1.5 Hz, 2 mm amplitude; C: 0.2 Hz, 2 mm amplitude; E: 1.5 Hz, 0.2 mm amplitude; F: 3 Hz, 0.2 mm amplitude

Apparent from both Figure 2.5 and Table 2.5 is the accuracy of the TFDI approach in determining displacement when frequencies are within the working range of the accelerometers used (note that, as discussed previously, the working range for the accelerometers used was 0.5-1000 Hz). For both data sets with a 0.5 Hz frequency, the TFDI approach was unsuccessful in determining the amplitude of the vibrations. For only one of the two cases was the frequency detected but only with moderate success (run C - 0.5 Hz and 2 mm amplitude). This emphasizes the importance of understanding the working range for instruments used when employing the TFDI. Meanwhile, the TFDI very accurately calculated both the displacement and frequency for all other runs of data. It should be noted that the frequencies and amplitudes used in these calculations are in a similar range to those used in ex vivo experiments discussed later. Based on these results, it can be concluded that the TFDI approach allowed for an accurate calculation of displacement values from acceleration data provided the frequency content of the data of interest lies within the working range of the accelerometers used.

### 2.4 Isolation of Road Features of Interest

For the laboratory cycling experiments, two different road signatures of interest were sought for comparison. As stated previously, the hexapod data collection process used three road signatures from the occupational data collection – a smooth freeway segment, a rough freeway segment, and a city street segment. Though other road segments from the occupational data collection could have been used for the laboratory cycling experiments, determining the response of the torso to simulations would have relied solely on a mathematical model without validation from the sternum data from the hexapod data collection. Therefore, to allow for comparison with collected data, it was decided that the two road segments for the laboratory experiments be

amongst the three used as inputs for the hexapod. Amongst the three occupational inputs into the hexapod, the smooth and rough freeway segments (henceforth referred to as the "freeway" and "expansion joint" inputs, respectively) were selected because of the relative similarities in the speed at which the segments were traversed as well as the differences in the vibrations imparted to the drivers as a result of the expansion joints. Therefore, the hexapod outputs for the freeway and expansion joint segments were selected for processing for use in the subsequent laboratory experiments with hopes of isolating a continuous exposure signature from the freeway segment and a more impulsive exposure signature from the expansion joint segment.

The freeway segment and the expansion joint segment both provided 180 seconds of data from which features and regions of interest needed to be selected for further processing. From the GPS data of the floor accelerations, two regions from the smooth freeway data set (25 and 12.5 seconds in length) were isolated which were traversed at a fairly constant speed with a consistent acceleration profile at the vehicle floor. The floor acceleration signals of these segments were also checked to ensure no acceleration values greater than 5  $m/s^2$  were present. In addition to this, four expansion joints (at about 9, 18, 50, and 90 seconds into the route) were isolated from the expansion joint data set. The location of these expansion joints were verified based on GPS data collected during the occupational data collection and the acceleration profiles of the expansion joints from the floor of the bus were checked to ensure no impulses greater than 5  $m/s^2$  were present within 1.5 seconds before and after the expansion joint of interest. The expansion joints were found to impart acceleration impulses greater than 5  $m/s^2$  while the regions of the freeway without expansion joints generally had acceleration values less than 5 m/s<sup>2</sup> The selected expansion joints were chosen such that the impulse imparted to the vehicle from the expansion joint and the subsequent oscillations reached pre-impulse levels prior to the vehicle encountering the next expansion joint. The six time frames (2 regions of the smooth freeway and 4 expansion joints) were thus selected as the features of interest to be used in the data processing for the laboratory experiments.

### 2.5 Mathematical Model of Vibration Response

#### 2.5.1 Description

As stated previously, because sternum acceleration data of subjects exposed to WBV was not initially available for use, a mathematical model was developed to allow for determining sternum responses to seat accelerations. Although sternum data later became available for use, the mathematical model was nevertheless developed to predict and verify the results that would eventually be extracted from the shaker table data collection. The vibration response of the seat-sternum system was mathematically modeled using a two-parameter mass-spring-damper mechanical model consisting of two components – one component modeling the vibration of the buttocks of the seated subjects and the other component to modeling the vibration of spine from the sacrum to the sternum. Both components consisted of a parallel mass-spring-damper system as shown in Figure 2.6 but with differing spring and damping coefficients, as determined from literature <sup>63, 64</sup>. The mathematical derivation of the model follows.



Figure 2.6 Mass-spring-damper mechanical model. M, Z, F, d, and k represent the mass, vertical displacement, input force, damping coefficient, and spring coefficient of the system, respectively.
If the forces related to the mass, dashpot, and spring are  $F_m = Mz$ ",  $F_d = dz$ , and  $F_s = kz$ , respectively, where single and double apostrophes indicate first and second derivatives of the position (z). Then, the sum of forces can be expressed as:

$$\sum F_{total} = F_m + F_d + F_s = F$$
$$\sum F_{total} = Mz'' - dz' - kz = F$$

Dividing the above terms by mass to get F/M (acceleration) gives:

$$z'' - \frac{d}{M}z' - \frac{k}{M}z = F/M$$

Substituting for damping ratio ( $\zeta$ ) and natural frequency ( $\omega$ ) with

$$\omega = \sqrt{\frac{k}{M}}$$
 and  $\varsigma = \frac{d}{2M\sqrt{kM}}$  then yields:  
 $z'' + 2\varsigma\omega z' + \omega^2 z = \frac{F}{M}$  (eq. 3)

# 2.5.2 Model Results

The differential equations associated with eq. 3 were then used with mathematical computing software (Matlab R2009, Mathworks, Natick, MA) to predict the response of the buttocks and the segment of the spine between the thighs and sternum to the floor accelerations. Seat pan acceleration data from the two vibration signatures for each subject was inputted into a differential equation solver which predicted the displacement response. This was performed to model both the buttocks and the individual intervertebral disc response to the input acceleration. Model parameters for the buttocks and spine model are given in Table 2.6. For the spine, the

natural frequency was calculated by summing the spring constant of the intervertebral discs across 7 equivalent segments. 7 equivalent segments were used because from sacrum to the sternum, 8-10 intervertebral discs are present, of which 5 are lumbar discs while the remainder are from the thoracic region of the spine. Compared to the thoracic intervertebral discs, lumbar discs are larger and better suited for bearing loads <sup>30</sup>. The larger size of the lumbar intervertebral discs enables greater tensile and compressive displacement compared to the thoracic intervertebral discs. Additionally, ribs are also present on all thoracic vertebrae, many of which are connected to adjacent ribs via cartilage linkages. These ribs further decrease the mobility of thoracic intervertebral discs. Therefore, to account for the decreasing mobility of intervertebral discs as one moved up the spine, the displacement values were divided across 7 equal segments. Displacement simulations were conducted for each of the 12 hexapod subjects for 180 seconds of seat acceleration data from both the freeway and expansion joint road segments.

	Stiffness (k)	Damping (d)	Damping Ratio (ζ)	Natural Frequency (ω)
Buttocks <sup>64</sup>	8500 N/m	35 Ns/m	0.01	0.624 rad/s (3.92 Hz)
Intervertebral Discs <sup>63</sup>	50000 N/m	Only ζ used	0.15	0.572 rad/s (3.59 Hz)

Once displacement data had been calculated for both the buttocks and the intervertebral discs from the simulations, the data needed to be processed to extract the amplitude of the displacements. For each of the four expansion joints, the mean peak-to-peak was calculated displacement for each subject and then an average displacement was calculated for the 12

subjects. For the freeway segments of interest, twice the root mean square displacement was calculated and averaged across the 12 subjects. The results of these calculations for both the buttocks and the intervertebral discs are presented in Table 2.7. Of note is the much larger displacement amplitude seen for the expansion joints compared to the freeway segments.

Table 2.7	Mathematical	model	results

	Freeway Segments (RMS)	Expansion Joints (peak)
Buttocks	1.44 mm	1.58 cm
Intervertebral disc	0.25 mm	2.18 mm

## 2.6 Analytical Results

Once a mathematical model had been developed, the data from the hexapod was processed to extract the response of the spine to the input vibrations. The acceleration data from both the seat and the sternum was first converted to displacement data using the TFDI approach. After this, the same 6 regions of interest (2 freeway segments and 4 expansion joints) discussed in section 2.3 were isolated from the displacement data for the seat and sternum. Local minima and maxima for the displacement data were then found using the peak detection algorithm discussed in section 2.3.2 for each expansion joint and each driver at the sternum and seat (96 waveforms total – 48 at the sternum and 48 at the seat). Once the local minima and maxima (peaks) of each of the four expansion joints had been detected for each subject, a composite waveform was created for each of the expansion joints by averaging the time and amplitude of the peaks at both the seat and the sternum across all 12 subjects. From four these composite expansion joint waveforms for the seat and the sternum, the peak-to-peak amplitudes from each local minima to local maxima were determined.

Similarly, the duration in time from each local maxima to local minima and from each local minima to local maxima were calculated. The average of these values at the seat and at the sternum were then calculated across all subjects and used to calculate the expansion joint vibration signatures for both the seat and the sternum presented in Table 2.8. Note the similarities in the frequency response of the seat and sternum.

Location	Frequency	Amplitude
Seat	2.73 Hz	9.4 mm
Sternum	2.71 Hz	36.3 mm

 Table 2.8
 Expansion joint seat to sternum signatures

The vibration signatures for the freeway segments were calculated using a similar protocol. First, the acceleration data at the seat and at the sternum for each subject and each of the two freeway regions of interest was converted to displacement data using the TFDI approach. The displacement data was then filtered using an 8<sup>th</sup> order dual-pass Butterworth low-pass filter (4<sup>th</sup> order each pass) with a cutoff of 30 Hz to remove high frequency noise from subsequent calculations. The 30 Hz cutoff was chosen based on ongoing spectral analysis work that has thus far indicated that most energy from WBV vehicle signals is between 0-15 Hz with little above 30 Hz in the direction perpendicular to the vehicle floor. After filtering, the peak detection algorithm discussed in section 2.3.2 was used to detect local minima and maxima at both the seat and sternum for each of the subjects and for each of the two freeway regions (450 total seconds of freeway data). The mean of twice the root mean square of the displacement amplitude values and the average time between successive peaks in the signal were then used to find the frequency

and amplitude of the freeway vibration signatures. The results are detailed in Table 2.9. Note the similar frequency and the smaller amplitude compared to the expansion joints.

Location	Frequency	Amplitude
Seat	2.43 Hz	2.6 mm
Sternum	2.59 Hz	6.4 mm

Table 2.9Freeway seat to sternum signatures

Once the freeway and expansion joint signatures had been calculated, signatures to be during the experimental cycling needed to be determined. First, the values of the displacement of the glutes/buttocks from the mathematical model were subtracted from the analytical results for the freeway and expansion joints. This calculation provided the displacement of the spine from the sacrum to the sternum. These results were then divided by 7 to again account for the anatomic variability of the spine and to give the displacement signatures of the intervertebral discs. Because the freeway and expansion joint segments had relatively similar frequencies of the two road types, a 2.5 Hz frequency was used for both signatures. The resulting values as well as predictions from the mathematical model are provided in Table 2.10.

Location	Model	Frequency	Amplitude
Frankov	Analytical 2.5 Hz		1.60 mm
rieeway	Mathematical	-	2.18 mm
Europeien Leinte	Analytical	2.5 Hz	0.34 mm
Expansion Joints	Mathematical	-	0.25 mm

Table 2.10Signatures used at start of preliminary tests

In general, the mathematical model provided displacement amplitude values that were similar to those extracted from the experimental model. Both indicated an amplification of the seat accelerations at the sternum and both indicated much higher displacement values for the expansion joints than the freeway segments. Due to the desire to use data that was as representative of occupational exposures as possible, the analytical model results were used as the input signatures for the experimental cycling. The degree to which these values were modified/altered for the final round of experiments is discussed in the next chapter.

### **CHAPTER 3: Preliminary Study and Protocol Refinement**

### 3.1 Preliminary Study Overview

A preliminary laboratory study was conducted prior to the final laboratory study. The purpose of the preliminary study was to finalize a laboratory protocol for loading human cadaver functional spine units (FSUs) with displacement signatures representative of continuous and impulsive occupational WBV exposures. As stated previously, this project was the first of its kind to use occupational WBV exposures in order to study spinal loading biomechanics in a laboratory setting. As such, protocols and procedures for doing so were nonexistent at the start of the project and needed to be developed. Prior to the start of the project, it was understood that developing and perfecting these protocols would require multiple iterations of specimen testing. Therefore, the purpose of the preliminary study was to use suboptimal human cadaver FSUs to iteratively develop the testing protocol prior to conducting the final laboratory tests.

Within the realm of perfecting the final testing protocol, the preliminary study had five main aims. These aims are as follow:

- 1. Outline final cycling signatures in terms of frequency, displacement, and number of cycles. Although processing signals from the shaker table data collection did provide frequencies and displacements of interest, the response of human tissues to these signatures still needed to be tested. Also, the number of cycles for each signature had to be determined.
- 2. *Develop and test a method by which disc damage could be traced.* One of the aims of this project was to develop a method of outlining disc damage which did not rely on radiography. This method needed to be tested prior to use in the final experiments.

- 3. Determine what types of damage occurs in the intervertebral discs and vertebral bodies *as a result of the cycling protocol.* Prior studies examining disc loading and/or cycling had reported migration of the nucleus, deterioration of the annulus, and/or endplate fracture <sup>31, 58, 59</sup>. It was not known what types of damage, if any, would occur in the intervertebral discs and vertebral bodies as a result of the extended cycling procedures.
- 4. Outline progression of disc damage. As stated previously, it was not known whether damage to the disc, if present, progressed from the interior of the intervertebral disc outward or from the exterior of the disc inward. The injury tracing protocol was developed under the assumption that damage progressed from the region of highest to lowest pressure (outward from the interior of the disc). It remained to be seen whether disc damage could be traced when the FSUs were subjected to extended continuous and impulsive cycling.
- 5. Determine whether FSUs from older human cadavers could be used. The cadaveric disc specimens to be used came from a tissue bank. For the most part, these disc specimens were from elderly donors and, as a result of gradual deterioration throughout the donor's lifetime, many of the vertebrae and intervertebral discs were not in their healthiest state. The injury tracing technique relied on the disc being intact and able maintain a nominal level of intradiscal pressure. The ability of the tracing technique to outline damage in discs which did not maintain pressure needed to be examined.

# 3.2 Specimen Preparation

For both the preliminary tests and the final experiments, whole human lumbar spine segments (T1-S5) were received frozen from a tissue bank (ScienceCare Inc; Phoenix, AZ) and stored in a freezer at -20° C. Functional spinal units (FSUs) of interest were extracted from the

spine segments by cutting through the intervertebral discs superior and inferior to the FSU of interest. All remnants of these sliced intervertebral discs were then removed from the endplates of the FSU of interest. Additionally, all ligamentous soft tissue, musculature, and bony structures posterior to the pedicles of the vertebral body were removed from the FSUs using a scalpel and bone saw, respectively. All extracted FSUs were dried, wrapped in a cloth, and stored in a freezer at -20° C at all times during which they were not being prepared or tested; FSUs were thawed prior to use by setting in a warm water bath for approximately 30 minutes.

Seven FSUs from 5 different spines were harvested for the preliminary study and refinement of the protocol for the final experiments. The integrity of each intervertebral disc was assessed by injecting an aqueous saline solution of 0.90% mass concentration of sodium chloride into the nucleus pulposus using a 10 ml pressure inflator (Model 710-111 Leveen; Boston Scientific; Boston, MA) and a 20 gauge needle. The injection site of the needle was marked on the exterior of the intervertebral disc using a semipermanent marker. A disc was deemed intact if it maintained an intradiscal pressure above 90 PSI with an injection of at most 3 ml of solution. The 90 PSI value is in line with the high-pressure endpoint of 100 PSI used during discography <sup>65, 66</sup>. The intradiscal pressure of each disc was monitored using the pressure inflator gauge for 2-3 minutes to determine if a drop in pressure of more than 10 PSI occurs. Henceforth, discs will be described as being either "intact" or "not intact" based on their ability to sufficiently maintain intradiscal pressure above 90 PSI in the test described. Donor details and information on the integrity of intervertebral disc used are provided in Table 3.1 on the next page.

After checking disc integrity, three metal screws were inserted through the cranial half of the superior-most vertebral body and the caudal half of the inferior-most vertebral body of each FSU. The screws were used to help stabilize the FSU in the PMMA mold. Henceforth, the cranial half of the superior-most vertebral body and the caudal half of the inferior-most vertebral body of each FSU will be referred to as "the potted region" of the FSU. The potted regions of the FSUs were then embedded in polymethylmethacrylate (PMMA) such that the top and bottom of each PMMA mold was near-parallel with the superior- and inferior-most endplates of the FSU. An image of an embedded FSU is shown in Figure 3.1.

Specimen ID#	Gender	Age at Death	Segments Used	Disc Intact?
180	Male	27 years	T11-T12	Yes
180	Male	27 years	L1-L2	Yes
500	Male	49 years	T12-L1	Yes
907	Female	66 years	T12-L1	Yes
056	Male	72 years	L2-L3	No
056	Male	72 years	L4-L5	No
044	Female	61 years	L2-L3	No

Table 3.1Specimens used in preliminary studies



Figure 3.1 Embedded FSU as used in tests. View shows anterior side of disc. Pink and blue colors on exterior of disc are from staining protocol outlined in section 3.2. Metallic structure above FSU is actuator of servohydraulic testing system.

# 3.3 Stain Injection Protocol

Prior to performing the preliminary experiments, a two-part ink staining technique needed to be developed to trace disc damage. The stains served two distinct purposes; one to establish the preexisting condition of the disc and the second to trace damage imparted by the exposure protocols. The first staining protocol was specifically designed to highlight the preexisting location of the intervertebral disc's nucleus pulposus and the preexisting condition of the inner annulus fibrosis. The second stain was selected such that it could elucidate any new damage to the disc or endplates while not masking the initial stain. Note that, as stated previously, an assumption was made that intervertebral disc damage would progress from the interior of the discs towards the exterior. As such, the developed technique sought to target features that were in the interior of the annulus towards the interior of the disc. The technique developed does not rely upon radiography-based inks or stains and provides a more efficient and potentially cost-effective alternative to x-ray-based detection of nuclear material, as used in previous studies <sup>58-60</sup>.

### 3.3.1 Stain Selection

Many different possible stains were identified and tested for their ability to achieve the desired experimental effects and a combination of two stains was found that met the testing requirements. These two stains were a 1% aqueous alcian blue solution (Electron Microscopy Sciences; Hatfield, PA) and a 2.5% aqueous safranin-O solution (Sigma-Aldrich Corporation; Saint Louis, MO). The selection of these stains was based predominantly upon a literature review of prior staining techniques used to highlight features of the intervertebral disc, but not the migration of the nucleus pulposus <sup>67, 68</sup>. However the targets of each of the stains allowed for a

combination of the two to be used with great effect for highlighting the before- and after-state of the intervertebral discs used in the experiment. Safranin-O has been shown as an effective stain for highlighting both collagen (as found in the inner annulus) and glycoproteins (as found in the nucleus pulposus), staining both a bright red/pink color <sup>69, 70</sup>. Conversely, the alcian blue stains glycoproteins, but not collagen, a bright blue color <sup>71</sup>. However, when the two stains are used sequentially, collagen remains a red color and proteoglycans turn deep purple in color. Figure 3.2 demonstrates the effect of these stains by showing the same intervertebral disc sliced in thirds and then stained with the safranin-O stain, the alcian blue stain, and sequentially using first safranin-O stain followed by alcian blue stain. Note how safranin-O stains collagen and nuclear material a red/pink color, alcian blue only stains nuclear material a blue color, and using both sequentially results in collagen remaining red and nuclear material turning deep purple in color. Thus, using two different stains in a sequential manner allowed for the coloring of the inner annulus of the disc red (via the safranin-O stain) to trace preexisting damage to the annulus fibers. The nucleus pulposus is first colored red (via the safranin-O stain) and is then turned a blue/purple color (via the alcian blue) after being stained by both stains. This blue/purple allows for the nucleus to be contrasted with the red color of the annulus fibers.



Figure 3.2 Before (left) and after (right) images of intervertebral disc stained three times. On right, clockwise from bottom: area stained with safranin-O stain, area stained with alcian blue stain, and area stained first with safranin-O and then with alcian blue stain. Approximate area of nuclear pulposus circled in yellow.

## 3.3.2 Stain Injections

Based on the ability of the stains used to target specific components of the intervertebral disc, the following staining protocol was developed for highlighting the pre- and post-experiment condition of the interior of the intervertebral disc. All injections were performed using a standard 10 ml plastic syringe and a 20 gauge needle using solely thumb pressure. Provided the intervertebral disc is able to sufficiently able to maintain intradiscal pressure during injections, all volumes of stain injected are with maximal thumb pressure. If the disc was unable to maintain sufficient intradiscal pressure during injections, the stain would follow the path of least resistance and flow outwardly towards the outer annulus rather than into the inner annulus and nucleus pulposus. If this is the case, the inner annulus will remain unstained and damage to the disc brought on during cycling cannot be adequately detected. This idea that the intervertebral discs must maintain sufficient intradiscal pressure during injections was tested by using not intact (as determined by the disc's inability to maintain intradiscal pressure above 90 PSI) discs during the preliminary tests. An overview of the staining protocol and the anticipated results from each step is provided below.

- Inject safranin-O stain into nucleus of intervertebral disc. This will outline the location of the inner annulus while also seeping into any preexisting fissures of the collagen fibers that may be present. Additionally, the safranin-O stain will also stain the nucleus pulposus a preliminary red/pink color. It is assumed the safranin-O will remain in the inner annulus for a sufficient period to stain the targets.
- 2. Allow time for the safranin-O stain to settle while also exposing the disc to low forces to allow for collagen fibers to stretch and the stain to permeate to maximally stain target areas.

This part of the staining protocol was introduced while testing the fourth of the preliminary test specimens, as detailed later in this writing.

- 3. Inject alcian blue stain into nucleus of intervertebral disc prior to exposure of FSU to the desired vibration signatures. This will target solely the nuclear material, leaving the collagen fibers red/pink from the safranin-O stain. When the alcian blue stain mixes with pre-existing safranin-O, the nuclear material will turn deep purple in color. If the volume of alcian blue is less than that of the safranin-O, the nucleus pulposus will take a light purple color, an effect which can be rectified in future steps by injection of more alcian blue stain. If the volume of the alcian blue is greater than that of the safranin-O, the nucleus will appear much darker in color. Once the nucleus has been saturated with stain, any excess stain from the injections will remain in the inner annulus and follow a path of least resistance from there.
- 4. Periodically, throughout the experiment, inject additional volumes of alcian blue stain into the intervertebral disc nucleus to account for degeneration of the disc and a resulting increased capacity of the intervertebral disc to hold volume.
- 5. The staining procedure allows for the identification of components of interest in the following manner: all collagen fibers of the inner annulus and all preexisting fissures will be a red/pink color; the nuclear material will be a deep purple color; any damage to the inner annulus and any migration of the nucleus pulposus will be highlighted by the presence of a bright blue or deep purple color without any red/pink present.

# 3.4 Overall Testing Protocol

For both the preliminary tests and the final experiments, the same general procedure was used with only variations in the disc loading procedures between the preliminary and final tests. Frozen, potted FSUs were thawed and first injected with the safranin-O stain. FSUs were then placed in a warm water bath for between 5-20 minutes after which they were dried and placed on the servohydraulic testing system (henceforth referred to as the "MTS"). The FSUs were centered under the actuator in both the sagittal and coronal plane. The specimen was placed on the MTS such that the direction of the MTS actuator's movement was perpendicular to the FSU endplates and the actuator would first contact the superior-most of the two vertebral bodies. The alcian blue was injected into the intervertebral disc prior to the start of cycling and midway through the cycling protocol. After cycling, the intervertebral disc was dissected along the superior-most endplate using a scapula. The disc was then sectioned into slices 1-2 mm in thickness from the superior to the inferior region of the disc.

The vibration signatures used during the preliminary tests were under modification throughout the preliminary study. As such, the preliminary study will be grouped into 3 phases, each with variations in how specimens were cycled. Specifics of the cycling protocols and the manner in which they were refined are detailed next.

# 3.5 Preliminary Study Inputs, Results, and Refinements

### 3.5.1 Phase 1 of Preliminary Study

For the first phase of the preliminary study, 4 specimens were used. 1 specimen was exposed to the freeway cycling signature and the other 3 were exposed to the expansion joint signature. All specimens were placed under a static preload of 400 N during cycling to correspond approximately to the upper body mass of an average US adult <sup>30, 72</sup>. However, the viscoelastic properties of the disc and the gradual relaxation of the disc annulus were not accounted for by this preload. All 4 specimens were cycled for 14400 cycles of the signature of

interest. The number of cycles was determined by the median of cycles used to induce prolapse in one disc cycling study and the maximum reported number of cycles for another disc cycling study, both of which resulted in the same number <sup>58, 60</sup>. For this preliminary test, after injecting the safranin-O stain, the FSU was not cycled prior to the addition of the first round of the alcian blue stain. The safranin-O stain was injected, the specimen was placed in a warm water bath for 20 minutes, and the alcian blue stain was injected thereafter. The zero displacement position of the MTS actuator was set to the location which induced the preload at the start of cycling. This point was used as the zero displacement reference point for the remainder of the cycling and was not readjusted. The FSU was then cycled for 7200 cycles after which it was re-injected with the alcian blue stain. This reinjection was to allow for the stain to account for the increased ability of the intervertebral disc to hold volume as its interior degraded. The specimen was then repositioned on the MTS and the zero displacement position of the actuator was set to that which had induced a 400 N preload at the start of cycling. The FSU was then cycled for an additional 7200 cycles and dissected. All dissections proceeded from the superior region of the intervertebral disc towards the inferior region of the disc by slicing the disc into 1-2 mm thick slices along the transverse plane.

The freeway and expansion joint cycling waveforms used for phase 1 of the preliminary tests are shown across a 4-second window in Figure 3.3. Note that the negative displacement refers to the distance travelled by the actuator in the downward direction after being in contact with the FSU at its superior-most point. The 0 mm displacement point in these and subsequent figures refers to the point at which the MTS actuator induces the desired preload to the specimen.



Figure 3.3 Cycling waveforms for the freeway signature (top, blue) and the expansion joint signature (bottom, red) for phase 1 of the preliminary study. Note how both waveforms were at a frequency of 2.5 Hz. Negative displacement represents distance travelled by the MTS actuator from the superior towards the inferior of the intervertebral disc.

The first of the preliminary study specimens (specimen ID# 500, T12-L1) was exposed to the freeway signature and showed no signs of deterioration upon dissection. The specimen was cycled using a peak-to-peak amplitude of 0.34 mm and a cycling frequency of 2.5 Hz. Upon dissection, no migration of nuclear material, no fraying of the annulus fibrosis, and no endplate fracture was detected. The nuclear material was deep purple upon dissection and remained within the outline of the location of the inner annulus as indicated by the safranin-O stain. An image from the first preliminary test specimen is shown in Figure 3.4. Please note that because this was among the first pictures taken of a specimen dissection for this experiment, the degree to which photographs could highlight results was not understood. Additionally, how stray collagen and nuclear material would impact the results indicated by the photographs was not well understood. As such, the disc was not cleaned of stray collagen and nuclear material prior to taking photographs. In the image shown, the red/pink region of the inner annulus is well defined for the superior side of the disc (on left). On the inferior side of the disc (right), the nucleus was stained a deep purple color and contained within the safranin-O outline. Stray collagen, nuclear material, and dye remnants from the scalpel are seen as the brighter blue areas of the image.



Figure 3.4 Results from first preliminary test. Images show the central region of the disc in the transverse plane. T12 shown on the left and L1 shown on the right.

The second preliminary test specimen (specimen ID# 907, T12-L1) was subjected to the expansion joint vibration signatures with the peak-to-peak amplitude and frequency of the vibrations being 1.5 mm and 2.5 Hz, respectively. 1.5 mm was used as the displacement instead of the 1.6 mm calculated from the hexapod shaker data – a fact later corrected during the final study upon review of the results from the data processing. A cycling protocol identical to that of the first preliminary specimen was followed apart from the change of the amplitude of the displacement (1.5 mm peak-to-peak instead of 0.34 mm). The disc was cycled for the same number of cycles, at the same frequency, and followed the same injection protocol.

Upon dissection of the disc, damage was detected to both the intervertebral disc and one of the vertebral bodies. Small amounts of migration of the nucleus pulposus beyond the outline of the safranin-O outline were found. Additionally, an endplate fracture was found on the superior endplate of L1 (inferior of the two endplates surrounding the intervertebral disc). The idea that this fracture was brought on during cycling was supported by the fact that prior to testing and upon injecting the safranin-O and the first round of alcian blue, the disc only took a limited volume of fluid (approx. 0.4 ml total) via maximum thumb pressure. After the first 7200 cycles, however, the disc was freely accepting the alcian blue stain (approx. 1 ml was injected). This lack of ability to maintain pressure is indicative of disc damage, as discussed in section 3.2. The presence of the endplate fracture was verified by finding a region of the superior L1 endplate having been pushed into the cancellous bone of the vertebral body. This was further verified by cutting both of the FSU's vertebrae in half transversely using a bone saw. This revealed a brownish-red color to the inside of T12, as expected of cancellous vertebral bone. The inside of L1, however, was a blue-green color as brought about by leaking of stain into the cancellous bone. This contrast in color is shown in Figure 3.5.



Figure 3.5 Results from second preliminary test. Note the blue-green color of the cancellous bone on the left. L1 shown on the left and T12 shown on the right. The difference in color is brought on by leaking of stain from the intervertebral disc, through the endplate fracture, and into the cancellous bone of the vertebra.

The third preliminary test specimen (specimen ID# 180, L1-L2) was subjected to the same vibration protocol as the second preliminary test specimen. The specimen was an intact disc that maintained intradiscal pressure (as defined by the ability to only accept a limited volume of fluid via injection with thumb pressure) during the safranin-O and initial alcian blue injections. The specimen did not, however, maintain pressure after the first 7200 cycles.

Dissection of the disc revealed small amounts of nuclear material beyond the outlines of the safranin-O stain, as can be expected of damage to the annulus fibers. Dissection of the FSU also revealed cracks and an endplate fracture along the superior L2 endplate (inferior of the two endplates adjacent to the intervertebral disc). The endplate fracture was verified by cutting the vertebra in the transverse plane using a bone saw, as with the second preliminary test specimen. The cracks were highlighted by alcian blue stain and radiated out from the center of the endplate. The endplate fracture and cracks are shown in Figure 3.6.

The location and progression of the cracks along the endplate helped shed some light on the validity of the assumption of outwardly progressing disc injury. The location of the cracks and their progression away from the center of the endplate appear to suggest damage that radiates away from the center of the disc. Because the damage in this case is along the hardboned endplate and not the intervertebral disc, the location and progression of these cracks cannot necessarily indicate the progression of injury in the intervertebral disc. However, they do indicate that the greatest forces and pressures along the endplate are experienced at the center of the endplate, as evidenced by the fact that the endplate fractures encountered in both cases thus far had been at the center of the disc. From this it seems reasonable that the greatest forces and pressures experienced by the intervertebral disc which is between the source of the force (the MTS actuator) and the endplate are also at or near the center of the disc. This is in accordance with the assumption made for tracing disc injury – that progression of disc injury was from the interior of the disc towards the exterior (outward from the region of greatest intradiscal pressure).



Figure 3.6 Image from third preliminary test. Image shows the anterior half of the superior L2 endplate. Note the ability of the alcian blue stain to trace endplate cracks as indicated by the blue arrows. The cracks appear to progress outwards from the center of the disc.

The fourth preliminary test specimen (specimen ID# 180, T11-T12) was subjected to the same impulsive vibration exposure as the second and third preliminary test. The protocol used to subject the FSU to the vibration exposures was identical to that used for the second and third preliminary tests apart from one modification. Between the injection of the safranin-O stain and the first round of alcian stain, the disc was cycled 10 times using the smooth freeway input (0.34 mm peak-to-peak displacement at 2.5 Hz) under a static preload of 400 N. This was to allow for the annulus fibrosis to stretch and contract, thereby preconditioning the disc and allowing greater penetration of the safranin-O stain as well as better highlighting of the preexisting condition of the interior of the disc. After these 10 cycles, the disc was injected with alcian blue stain and an identical procedure as that for the second and third preliminary test specimens was resumed.

The specimen used was again an intact disc that maintained intradiscal pressure during the safranin-O and initial alcian blue injections. After the first 7200 cycles, the disc no longer

maintained pressure when injected with a large volume of alcian stain (over 1.0 ml). This is indicative of damage induced by the cycling protocol. Dissection of the disc revealed a greater initial inner annulus area than previously seen. This can be attributed to the cycling between the safranin-O and first alcian blue injections. There was no migration of the nucleus pulposus or fraying of the inner annulus. However, the nuclear material did seem to have sunk towards the inferior of the intervertebral disc as shown by the image of disc in Figure 3.7 on the next page. This sinking of the nuclear material along the cranial-caudal axis is likely attributable to the fact that there was an endplate fracture on the superior endplate of the inferior vertebra, as encountered with all previous preliminary specimens exposed to the rough freeway exposure. This was again verified by observing the color of the cancellous bone after cutting the vertebrae in half along the transverse plane.

Beyond the endplate fracture, the test specimen also highlighted the ability of the staining protocol to identify preexisting fissures in the annulus of the disc. A large fissure was found protruding from the center of the nucleus pulposus towards the anterior end of the disc. Alcian blue was found to have seeped into the fissure. However, the fissure was determined to have existed prior to cycling because of a bright red/pink outline surrounding the alcian blue stain. This red outline was indicative of the safranin-O stain penetrating the fissure prior to the alcian blue stain injections. It should be noted that despite the presence of this fissure, the disc was still able to maintain sufficient intradiscal pressure to be deemed intact prior to cycling. This would indicate that the fissure did not penetrate beyond the exterior of the annulus fibrosis and provide support to the idea that WBV-related disc injuries do progress from the interior of the interior of the interior of the exterior and not the reverse. Had the injury progressed in the reverse direct, it is likely the exterior of the annulus would have been ruptured so as to allow

stain to escape. This ability to highlight preexisting damage to the inner annulus was also a requirement of any staining technique that was to be used for the final cycling experiments. Images of the fissure and the stain outlines are shown in Figure 3.7 on the next page.



Figure 3.7 Fissure found in fourth preliminary test. Image on left shows central region of intervertebral disc above T12. Anterior end of the disc is towards the bottom of the image. Image on right is a magnified copy of the area outlined in blue in image on left. Note the red/pink outline surrounding the blue ink in the fissure. Blue ink on upper right of image on left indicates injection site.

Results from phase 1 of the preliminary study indicated that, with similar exposure variables apart from displacement amplitude, the expansion joint signature was more likely to induce damage to the discs than the freeway signature. The expansion joint exposure produced signs of annulus deterioration and endplate fractures while the freeway exposure produced no detected damage. The expansion joint signature causing more damage may be expected given the fact that only displacement varied between the two exposure profiles used.

Beyond this, phase 1 of the preliminary study also found that the disc staining technique capable of outlining intervertebral disc damage. The difference in color between the two stains was distinct to the point that differences between the inner annulus and the nucleus pulposus could be visibly recognized. That is, the contrast between the red safranin-O stain highlighting

the location of the inner annulus and the deep purple color of the mixture of the safranin-O and alcian blue stains highlighting the location of the nucleus pulposus allowed for easy detection of when the nuclear material extended beyond the area outlined by the initial safranin-O staining. Additionally, the staining technique also proved successful in tracing highlighting preexisting injury to the disc (as seen by the fissure in the fourth test specimen) as well as cracks along the vertebral endplates (as seen in the third test specimen). These results provided confidence in the ability of the staining technique to highlight damage of interest.

## 3.5.2 Phase 2 of Preliminary Study

Phase 2 of the pilot study sought to establish a framework to allow for a better comparison of the two exposures. Because both the smooth and rough freeway exposures had identical protocols outside of the amplitude of the vibrations, it was felt that the two exposures were essentially comparing different amplitudes of vibration without accounting for driver exposure to these different road conditions. It was decided to equate the two exposures based on drive time. For the freeway exposure, the frequency of the impulses imparted to the driver were no different than the rate at which the impulses were encountered (i.e. the road provided a continuous exposure to vibrations). For the expansion joint exposure, though the vibrations imparted an impulse at a frequency near 2.5 Hz, the expansion joints themselves were spaced about 50m apart and encountered at a frequency near 0.5 Hz (once every 2 seconds when travelling at a speed near 90 kmph). Thus, for an equivalent driving period, the number of impulses encountered during expansion joint driving was about one-fifth that encountered during the freeway exposure (0.5 Hz vs. 2.5 Hz). Therefore, it was decided that the rough freeway exposure should be modified such that the frequency of oscillation be 0.5 Hz to better

approximate temporal occurrence of the vibration exposure. The freeway and expansion joint cycling waveforms for phase 2 of the preliminary tests are shown in Figure 3.8. The 0 mm displacement point again represents the location of the MTS actuator at induction of preload.



Figure 3.8 Cycling waveforms for the freeway signature (top, blue) and the expansion joint signature (bottom, red) for phase 2 of the preliminary study. Note the difference in frequency between the two waveforms. Negative displacement represents distance travelled by the MTS actuator from the superior towards the inferior of the intervertebral disc.

For the second phase of the preliminary study, 2 specimens were used. 1 specimen was exposed to the freeway cycling signature and the other was exposed to the expansion joint signature. A few modifications were made to the cycling protocol from the first phase of the preliminary study. First, after the injection of safranin-O, the specimen was placed under a static 300 N load for 10 minutes to better allow for the disc to stress relax, as done by another study looking at migration of the nucleus pulposus <sup>59</sup>. All specimens were then placed under a static

preload of 200 N instead of 400 N because of the time allowed for the disc to stress relax. The specimens were then cycled 10 times using the freeway exposure signature to better allow for the safranin-O stain to highlight interior features of the intervertebral disc (this was identical to what was done for the fourth specimen of the first phase of the preliminary study). Then, the specimen was injected with the first round of alcian blue stain and the specimen was cycled for one-half the desired number of cycles using the exposure signature of interest. As stated previously, the number of cycles for the freeway exposure was 5 times that of the expansion joint exposure (14400 vs. 2880, respectively) to equate the drive times between the two signatures. After the first half of cycling, the specimen was then removed from the MTS, re-injected with alcian blue stain, and reset on the MTS. The zero displacement point for the MTS actuator again corresponded to the point at which the preload (in this case, 200 N) was induced during the initial one-half of the target number of cycles. The specimen was then removed from the MTS and dissected. It should be noted that both the specimens used for the second phase of the study were deemed to be not intact based on the discs' inability to sufficiently maintain intradiscal pressure. The discs were used to examine whether discs that were not in a completely intact state could be used during the cycling experiments. Because the specimens were unable to maintain a high intradiscal pressure at the start of the experiment, the maximum volume injected during any of single injection was 0.5 ml.

The fifth preliminary test specimen (specimen ID# 056, L2-L3) was exposed to the freeway cycling signature. Upon dissection of the disc, a large pool of deep purple stain was found in the posterior of the disc, behind the nucleus. This was a result of fraying of the inner annulus and deterioration of the annulus fibers. However, due to the large volume of stain injected into the disc and the fact that the disc was not intact at the start of the test, determining

whether this deterioration was the result of the cycling protocol was inconclusive. This was because the safranin-O stain did not adequately outline the location of the inner annulus during the initial injections. Due to the health of the disc and its inability to sufficiently maintain intradiscal pressure, the safranin-O stain did not localize at the nucleus pulposus but instead followed a path of least resistance away from the center of the disc. The alcian stain is believed to have then followed a similar path, thereby causing the pool of stain found upon dissection. No endplate fracture and no cracks were found along either of the endplates adjacent to the intervertebral disc.

The sixth preliminary test specimen (specimen ID# 044, L4-L5) was subjected to the modified expansion joint vibration protocol. The frequency of oscillation was set to 0.5 Hz and the number of cycles was set to one-fifth their previous value. The displacement amplitude was 1.5 mm peak-to-peak, as it had been for phase 1 of the preliminary study. Upon dissection of the disc, a large pool of deep purple stain was again found in the posterior region of disc. As was the case previously, determining whether this pool of stain was a result of the cycling procedure was inconclusive. No endplate fracture was detected on either of the endplates surrounding the disc and no fissures were found in the disc.

Though the modified expansion joint exposure better accounted for the road exposure time, the rate at which the vibrations occurred (2 seconds to compress and decompress the disc), was not like those encountered by bus drivers. The rate at which the impulses were encountered was correct at 0.5 Hz but the vibration signature of the impulses at 2.5 Hz was unaccounted for. As such, this loading protocol appeared to give the annulus fibers too much time between periods of contraction and relaxation for damage to the intervertebral disc to occur. A modification was made during phase 3 of the preliminary study to better account for the frequency differences.

The second phase of the pilot study highlighted the necessity to use intact discs when using the devised staining protocol. Both discs in this phase of the study were unable to sufficiently maintain intradiscal pressure and, as such, highlighting damage within the disc was unsuccessful. Both discs showed signs of deterioration to the inner annulus upon dissection but because no pressure existed to drive the stain towards the nucleus pulposus, the preexisting state of the inner annulus was not highlighted. Instead, the safranin-O stain pooled at a location outside the inner annulus and the alcian blue followed the same path during injections. As such, differentiating between preexisting damage to the discs and damage induced during disc cycling was without conclusive results.

# 3.5.3 Phase 3 of Preliminary Study

The vibration signature of the expansion joints was modified to account for both the frequency of the oscillations imparted by the impulses encountered by the bus drivers (2.5 Hz) as well as the rate at which these impulses were encountered (0.5 Hz). It was decided to maintain the vibration signatures at 2.5 Hz but introduce a delay between each cycle to account for the rate at which the impulsive exposures associated with the expansion joints were encountered (0.5 Hz). Additionally, to better account for the fact that the impulses imparted by the expansion joints were a decaying sinusoid, a second impulse of one-half magnitude of the first (0.75 mm) was introduced immediately after the first. Therefore, the cycling protocol for the expansion joint was modified such that each cycle contained three distinct phases: a 1.5 mm peak which occurred over 0.4 seconds, and a delay of 1.2 seconds. Note that cycle occurring in a time of 0.4 seconds corresponds to a frequency of 2.5 Hz.

was accounted for as was the frequency of the impulses imparted to the drivers by the expansion joints. The freeway exposure signature was kept the same as before and the number of cycles for each of the two exposure signatures was kept the same as for phase 2 of the preliminary study. Figure 3.9 shows the freeway and expansion joint cycling waveforms used during phase 3 of the preliminary study.



Figure 3.9 Cycling waveforms for the freeway signature (top, blue) and the expansion joint signature (bottom, red) for phase 3 of the preliminary study. Two cycles of the expansion joint exposure signature are shown in the diagram. The expansion joint waveform was designed to account for the frequency of the impulses (2.5 Hz) as well as the rate at which these impulses are encountered (0.5 Hz).

. Only 1 specimen was used during phase 3 of the preliminary study. This specimen was exposed to the modified expansion joint cycling protocol. All other aspects of disc cycling were identical to phase 2 of the preliminary study except the frequency and amplitude of the applied exposures. The disc specimen used was unable to sufficiently maintain intradiscal pressure.

The seventh preliminary test specimen (specimen ID# 056, L4-L5) was subjected to the modified expansion joint exposure signature. After dissecting the disc, a large pool of deep purple stain was again found in the posterior region of the disc. Determining whether this reservoir of stain was a result of cycling was inconclusive again due to the inability of the disc to maintain sufficient intradiscal pressure. Endplate fractures were, however, observed on both endplates adjacent to the intervertebral disc. Because the disc was unable to sufficiently maintain pressure at the start of the experiment, it could not be conclusively determined that the endplate fracture was brought on by testing. That said, it was felt that, amongst the expansion joint exposures used in the preliminary tests, this modified expansion joint signature most closely represented the occupational expansion joint exposure and the expansion joint signature devised for phase 3 of testing was used for the final cycling tests.

### 3.6 Preliminary Study Results

In all, the preliminary study provided answers to many of the questions posed at the start of phase 1 of the study. A review of some of the goals for the preliminary study as framed in section 3.1 follows.

- 1. *Outline final cycling signatures in terms of frequency, displacement, and number of cycles.* Each phase of the preliminary study outlined potential modifications to the cycling protocols. By the end of phase 3 of the preliminary study, cycling protocols for both the freeway and expansion joint exposures had been established.
- 2. *Develop and test a method by which disc damage could be traced.* The double-staining protocol using safranin-O and alcian blue stains was found to effectively trace and outline disc damage as brought on by cycling.

- 3. Determine what types of damage occurs in the intervertebral discs and vertebral bodies *as a result of the cycling protocol.* During the cycling tests, deterioration of the inner annulus, migration of the nuclear material, and endplate fractures were all encountered.
- 4. *Outline progression of disc damage.* This aspect of the preliminary testing goals remained inconclusive for the unhealthy discs used. For the healthy discs, not enough detectable damage was present to derive any conclusive results. Some evidence, however, did exist to indicate the location of greatest pressure during cycling is indeed at the center of the intervertebral disc (as assumed). Top increase chances of disc damage, the number of cycles used for the final tests were increased by a factor of 150%.
- 5. *Determine whether FSUs from older human cadavers could be used.* Based on the inability to determine the presence of disc-related injury in phases 2 and 3 of the preliminary study, it can be concluded that discs must sufficiently maintain intradiscal pressure to be used with the staining procedure. This is required for the safranin-O stain to reach the inner annulus and settle therein. When the disc could not maintain sufficient intradiscal pressure, pools of stain were found away from the center of disc.

An overview of the preliminary testing protocols is detailed in Table 3.2 and an overview of the results of the preliminary study is given in Table 3.3. Both Tables are on the next page.

Evident from the results of preliminary testing was that endplate fractures were only encountered in instances where the exposure was that of the expansion joints at frequencies of oscillation at 2.5 Hz. Also, apparent from the preliminary studies was the need to use healthy discs to effectively outline the state of the inner annulus prior to testing and prevent pooling of stain. Healthy discs were also required to effectively determine the extent of damage caused by the cycling protocol and whether endplate fractures were induced during testing or had existed prior to. The next chapter will detail the final round of experiments performed across six subjects using two standardized protocols.

Specimen	Segment	Exposure	Preload	Frequency	Amplitude	Cycles
500	T12-L1	Freeway	400 N	2.5 Hz	0.34 mm	14400
907	T12-L1	Exp. Joint	400 N	2.5 Hz	1.5 mm	14400
180	L1-L2	Exp. Joint	400 N	2.5 Hz	1.5 mm	14400
180	T11-T12	Exp. Joint	400 N	2.5 Hz	1.5 mm	14400
056	L2-L3	Freeway	200 N	2.5 Hz	0.34 mm	14400
044	L4-L5	Exp. Joint	200 N	0.5 Hz	1.5 mm	2880
056	L4-L5	Exp. Joint	200 N	2.5 / 0.5 Hz	1.5 / 0.75 mm	2880

Table 3.2Preliminary study protocols

Table 3.3Preliminary study results

Phase	Specimen	Segment	Intact?	Exposure	Annulus Damage?	Endplate Fracture?
1	500	T12-L1	~	Freeway		
1	907	T12-L1	~	Exp. Joint	~	~
1	180	L1-L2	$\checkmark$	Exp. Joint	$\checkmark$	$\checkmark$
1	180	T11-T12	$\checkmark$	Exp. Joint		$\checkmark$
2	056	L2-L3		Freeway	?	
2	044	L4-L5		Exp. Joint	?	
3	056	L4-L5		Exp. Joint	?	~

#### **CHAPTER 4: Final Study and Results**

## 4.1 Final Study Overview

The final round of tests had three primary objectives. These were as follow:

- 1. Determine whether either of the two vibration signatures was injurious to the spine. The preliminary tests were successful in outlining a protocol whereby occupational vibration exposures could be translated into a laboratory setting. What remained to be done was to compare the two exposures of interest and determine if either causes damage to the FSUs under the final protocol loading conditions.
- 2. Determine whether the same trends of disc and lumbar body degeneration observed in the preliminary study occurred under the final protocol loading conditions. In the preliminary study, the most definitive type of damage seen was endplate fractures. These endplate fractures were seen only amongst specimens exposed to the expansion joint signatures with impulses at 2.5 Hz. It remained to be seen if this was a consistent trend.
- 3. *Outline the progression of disc damage.* Due to not using all intact discs, the preliminary study provided little evidence as to whether disc damage progresses from the interior of the disc towards the exterior or in some other fashion. The final tests sought to provide a stronger understanding of this damage progression.

#### 4.2 Final Study Protocol

For the final tests, 6 FSUs with healthy intervertebral discs were used. Initially, the intent was to use 12 intervertebral discs for the final tests but, after initial pressure tests, only 6 were deemed to be intact and could sufficiently maintain intradiscal pressure. As discussed during the

preliminary tests, the staining protocol relied on intervertebral discs maintaining intradiscal pressure in order to indicate if disc damage was brought on as a result of cycling. As with the preliminary tests, discs were deemed intact if they were able to maintain an intradiscal pressure of 90 PSI with an injection of at most 3 ml of saline solution. Three of the 6 FSUs consisted of L2-L3 segments and three consisted of L4-L5 segments. Additionally, two pairs of FSUs were from the same donors. For the paired specimens, each of the two specimens was cycled using different exposures. Additionally, each segment for the one pair (be it L2-L3 or L4-L5) was cycled using a different exposure than the same segment for the other pair. Additional information on the specimens used is provided in Table 4.1.

Specimen ID#	Gender	Age at Death	Segments Used	Exposure
033	Male	65 years	L4-L5	Exp. Joint
044	Female	61 years	L2-L3	Freeway
067	Female	21 years	L2-L3	Exp. Joint
067	Female	21 years	L4-L5	Freeway
073	Male	54 years	L2-L3	Freeway
073	Male	54 years	L4-L5	Exp. Joint

Table 4.1Specimens used in final testing

The potting of specimens in PMMA and the storage thereof was identical to that of the preliminary tests. Each specimen was thawed in a warm water bath for approximately 30 minutes after which safranin-O was injected into the nucleus pulposus of the disc using a standard 10 ml syringe and a 20 gauge needle to elucidate the preexisting condition of the disc. As in the preliminary tests, all injections were performed by applying thumb pressure until maximum resistance was reached. At most 1.5 ml of stain was injected during a single injection.

As in the preliminary studies, the FSU was placed on the MTS and centered with the MTS actuator in both the sagittal and coronal planes. The FSU was placed on the MTS with the direction of the MTS actuator's movement perpendicular to the FSU endplates. Also, the FSU was set so the actuator would first contact the superior of the two vertebral bodies, as shown in Figure 4.1. The potted FSU was then placed under a 300 N static load using the MTS for 10 minutes to allow for stress relaxation of the annulus fibers. Then, the 300 N preload was reset and the disc was preconditioned by cycling10 times under the smooth freeway input signature (0.34 mm peak-to-peak displacement at 2.5 Hz). Following preconditioning, the specimen was removed from the MTS and the disc was injected with the alcian blue stain and repositioned in the MTS. The 300 N preload was reestablished and the position of the MTS actuator was set as the zero displacement position for the remainder of the test. The FSU then underwent three rounds of cycling using either the freeway input signature or the expansion joint input signature. The number of cycles for each round of cycling varied by input signature with FSUs exposed to the freeway input cycled 7200 times per round of cycling and those exposed to the expansion joint input cycled 1440 times per round of cycling. A flowchart of the cycling protocol is provided in Figure 4.2



Figure 4.1 MTS actuator in contact with specimen. MTS actuator is in contact with superior of the two vertebrae of the FSU



Figure 4.2 Flowchart of final testing protocol

After each round of cycling, the FSU was removed from the MTS and injected with another round of alcian blue stain. The FSU was then repositioned as before on the MTS and subjected to the next round of cycling. The zero displacement point of the MTS actuator was defined as the point where the preload was applied to the FSU at the start of cycling. This zero displacement point was not adjusted for any subsequent round of cycling. After three rounds of cycling and subsequent injections of alcian blue, the FSU was removed from the MTS and the intervertebral was dissected from the superior endplate towards the inferior using a scalpel by making slices 1-2 mm thick.

## 4.3 Final Test Results

Images from the dissection of each FSU are presented as a grid. The upper-left image of this grid shows the superior of the two endplates surrounding the intervertebral disc. The upperright image shows the superior region of the intervertebral disc as viewed from the superior
towards the inferior. The bottom-left image shows the transverse center of the intervertebral disc as viewed from the superior towards the inferior. The bottom-right image shows the inferior of the two endplates adjacent to the intervertebral disc. Figures 4.3 and 4.4 outline this image grid and the locations/views of each of the sections shown in the image grid.

Superior Endplate	Superior Region of		
Adjacent to Disc	Intervertebral Disc		
Center of	Inferior Endplate		
Intervertebral Disc	Adjacent to Disc		

Figure 4.3 Diagram of picture grids used to display results



Figure 4.4 Diagram of FSU (left) and picture grid (right). Lines through intervertebral disc on FSU correspond to location of images detailed in diagram of picture grid. Views shown in picture grids correspond to those written in each section of the diagram of grid.

## 4.3.1 Freeway Vibration Cycling Results

None of the three discs exposed to the freeway exposure signature showed any definitive signs of nucleus migration or fraying of the inner annulus fibers. Nor did any of these discs have an endplate fracture on either of the two endplates surrounding the intervertebral disc. What follows is a specimen-by-specimen presentation (case study) of the results of the intervertebral disc dissections.



Figure 4.5 Results from first test specimen (044, L2-L3) exposed to the freeway signature

Results from the first test specimen (specimen ID# 044, L2-L3) are shown in Figure 4.5. The disc maintained intradiscal pressure for all five injections (saline, safranin-O, and three alcian blue injections). Upon dissection of the disc, no damage was found to the superior endplate. The nuclear material was entirely within the outline of the safranin-O stain at the superior region of the intervertebral disc, as shown by the upper-right image of Figure 4.5. There was also no visible damage to the annulus fibers at the superior region of the intervertebral disc. Upon slicing further towards the inferior region, there appeared to be small volumes of the nuclear material extending slightly beyond the outline of the safranin-O stain and towards the

anterior region of the disc, however, the degree to which this nuclear material penetrated the boundary of the safranin-O stain was difficult to determine visually. The bottom-right image of Figure 4.5 demonstrates shows the degree to which the safranin-O stain outlined the location of the nucleus pulposus upon dissection. To the left of this image is the injection site of the disc, which is highlighted by parallel lines of stain extending from the exterior of the disc towards the nucleus pulposus. Upon reaching the inferior endplate, small amounts of both stains was found on tissues above the endplate. No endplate fracture was found on the inferior endplate.



Figure 4.6 Results from second test specimen (067, L4-L5) exposed to the freeway signature

The second test specimen (specimen ID# 067, L4-L5) exposed to the smooth freeway exposure was from very young donor (21 years old at death) and was therefore in very good condition at the start of the experiment. A specimen from the same donor was used as the second test specimen for the rough freeway exposure. This specimen also maintained intradiscal pressure during all five injections. Pictures from the dissection of this specimen are shown in Figure 4.6 in a manner and layout consistent with the diagram shown in Figure 4.3.

No discernible damage was found upon dissection of the disc. The upper-left image of Figure 4.6 shows the superior endplate adjacent to the intervertebral disc. No damage or deterioration of the endplate was found during dissection. The upper-right image shows the superior region of the intervertebral disc. Again, no damage was found during dissection and the nuclear material was found to be contained within the outline of the safranin-O stain. Slicing through the intervertebral disc towards the center of the disc also found no visible damage to the annulus fibers or migration of the nuclear material away from the center. Interestingly, the safranin-O stain indicated the disc had a rather non-symmetric inner annulus. As evidence of this, note the area stained by safranin-O in the upper-right image of Figure 4.6 versus that in the lower-left image of Figure 4.6. In both cases, the area of safranin-O stain is slightly to the left of the center of the intervertebral disc. However, going from the superior of the intervertebral disc towards the inferior of the intervertebral disc, the location of the inner annulus appears to shift from the posterior region of the disc towards the anterior. Despite this shift, the nuclear material was found to have been contained within the outlines of the safranin-O stain throughout the disc. Upon reaching the inferior of the two endplates, no damage or endplate fractures were detected.

The third specimen exposed to the freeway cycling protocol (specimen ID# 073, L2-L3) also showed no signs of deterioration as brought on by the cycling protocol. Results from this specimen are shown in Figure 4.7. The intervertebral disc again maintained intradiscal pressure during all five injections. Upon dissection, no damage was found to the superior of the two endplates, as shown in the upper-left image of Figure 4.7. The superior region of the intervertebral showed a very large area of inner annulus stained by the safranin-O stain relative to the amount of nuclear material present. This is shown in the upper-right image of Figure 4.7, particularly by the lack of nuclear material present towards the posterior region of the disc. In

previous dissections, this had been indicative of nuclear material sinking towards the inferior region of the disc. Slicing the intervertebral disc towards the inferior region revealed how much of the nuclear material appeared to have sunk towards the inferior region of the disc, as shown by the lower-left image of Figure 4.7. Whether this was a result of the testing or preexisting for this disc was inconclusive as the nuclear material remained within the outline of the safranin-O stain throughout the disc. Upon reaching the inferior of the two endplates, no endplate fracture was found. That said, the top of the endplate was chipped by the scalpel during dissection, thus introducing the cracks visible in the bottom-right image of Figure 4.7.



Figure 4.7 Results from third test specimen (073, L2-L3) exposed to the freeway signature

In all, the smooth freeway exposure resulted in no detectable damage to the intervertebral disc and no endplate fractures as a result of cycling. Considering the experiment mimicked about 2.4 hours (21600 cycles at 2.5 Hz) of driving exposure on a fairly smooth freeway surface at a relatively constant and attainable speed, the lack of damage to the intervertebral disc may not be all that surprising given the relatively short exposure duration. It should be noted that the model

used was a disc-body-disc model and therefore varies from anatomic exactness due to the lack of posterior vertebral structures and the lack of muscles and ligaments to support the FSU. Interpretation of the results from this disc-body-disc model are given later in this discussion but must be made with the caveat that variations from in vivo conditions are inevitable.

#### 4.3.2 Expansion Joint Vibration Cycling Results

Each of the three intervertebral discs exposed to the rough freeway cycling signature had an endplate fracture along the inferior of the two endplates adjacent to the disc. Additionally, two of the three discs also showed a deterioration of the inner annulus and/or migration of nuclear material beyond the outlines of the inner annulus, as colored by the safranin-O stain. What follows is a specimen-by-specimen analysis of the dissections of the disc specimens.



Figure 4.8 Results from first test specimen (033, L4-L5) exposed to the expansion joint signature

The first specimen (specimen ID# 033, L4-L5) exposed to the expansion joint signature had considerable preexisting and induced damage to the inner annulus. This is shown in the upper-right image of Figure 4.8. The safranin-O stain was able to outline an area towards the

anterior region of the disc where stain accumulated after entering at the injection site, which was at the posterior-right region of the image. Nevertheless, the disc did maintain intradiscal pressure during the saline injections, safranin-O injection, and the first of the alcian blue injections. The superior region of the intervertebral disc had alcian blue stain towards the posterior of the disc from the nucleus pulposus. Due to the color of the stain (blue, not deep purple) and the lack of an outline of safranin-O, the presence of the stain was indicative of damage induced by cycling procedure. The nucleus was located right of center and towards the posterior region of the disc. Additionally, some alcian blue stain gathered near the injection site at the superior of the disc and decreasing amounts were found in the same area as one moved towards the inferior region of the disc. The central region of the disc. This is shown in both the upper-right and lower-left images of Figure 4.8. The streaks of ink to the anterior left of the nucleus pulposus in the upper-right image is the same pool of ink seen in the anterior left of the lower-left image seeping through annulus fibers.

The inferior endplate of the specimen had a very large endplate fracture apparently brought about by the endplate caving at multiple sites near the center of the disc. The hard bone of the endplate was found to have been pushed about 1.5 mm into the cancellous bone of the vertebra. This is shown in the bottom-right image of Figure 4.8 with the large blue/purple region in the center of the vertebral body being the site of the endplate fracture. Additionally, cracks were also found along the inferior endplate, radiating towards the anterior region of the disc from the center. This is evidenced in the lower-right image of Figure 4.8 by the fissure radiating towards the anterior right of the disc from the center.

The second of the three specimens (specimen ID# 067, L2-L3) exposed to the expansion joint cycling protocol was from the same donor as the second specimen exposed to the freeway cycling protocol. The disc was from a relatively young donor and in good condition. The nuclear material remained within the outlines of the safranin-O stain throughout the disc, as evidenced by the upper-right and lower-left images of Figure 4.9. The injection site was at the left of the disc and stain remnants from injections can be seen in the lower-left image of Figure 4.9.



Figure 4.9 Results from second test (067, L2-L3) specimen exposed to the rough freeway signature

Despite the lack of any detectable damage to the inner annulus, an endplate fracture was found on the inferior endplate. This is visible on the lower-right image of Figure 4.9. Whereas endplate fractures in other specimens were characterized by a cracking of the vertebra's hard bone and its caving into the cancellous bone of the vertebra, the endplate fracture for this particular specimen showed no other signs of cracking or deterioration other than the fracture (visible as the deep blue line through the center of the vertebral body in the lower-right image of Figure 4.9). The fracture itself was slightly over a millimeter wide at its widest point and large enough to allow for ink to seep into the vertebral body. Interestingly, the disc maintained pressure for four of the five rounds of injections but did not hold pressure during the final alcian injection. This pattern of maintaining intradiscal pressure appears indicative of a gradual deterioration of the endplate whereas the fracture itself seems much more characteristic of what would be expected of a single event which did not impact other regions of the vertebral endplate.



Figure 4.10 Results from third test specimen (073, L4-L5) exposed to the expansion joint signature

The final test specimen (specimen ID# 073, L4-L5) exposed to the expansion joint cycling protocol was from the same donor as the third test specimen exposed to the smooth freeway cycling protocol. As was the case with the third specimen exposed to the freeway cycling protocol, considerable preexisting damage was outlined by the safranin-O stain, as can be seen throughout the images in Figure 4.10 (five images are shown as part of Figure: upper-left image shows the superior of the two endplates, upper-middle image shows the superior region of the intervertebral disc, upper-right image shows a more inferior region of the disc than the upper-middle image, lower-left image shows the inferior of the two endplates). The preexisting damage to the disc is most evident in the upper-right and lower-left images of Figure 4.10, as

there was a pooling of safranin-O stain in the left of the intervertebral disc. This is near the injection site, which was on the posterior-left region of the disc. That said, there was also a large amount of alcian blue stain which had migrated to the posterior-right region of the disc, as shown in the upper-middle and upper-right images of Figure 4.10. This was indicative of post-cycling damage to the disc as safranin-O stain was not found to have penetrated into the same area despite the intervertebral disc initially maintaining intradiscal pressure during the saline injection, the safranin-O injection, and the first of the alcian blue injections. Further alcian blue migration was found in the central region of the disc, where the stain migrated from the nucleus towards the anterior region of the disc, pooling in a crescent-like shape seen in the lower-left image of Figure 4.10. This same pool of ink can be seen seeping through the annulus fibers slightly anterior to the nucleus pulposus in the upper-right image of Figure 4.10.

The inferior endplate of the two endplates adjacent to the intervertebral disc had a large fracture very similar to those seen in other specimens, where a region of the hard bone of the vertebrae was found to have caved in. To demonstrate the degree to which this happened in this particular specimen, a ruler was used to help create the images shown in Figure 4.11. The ruler was inserted beneath the region of bone which had not been damaged and pushed towards the posterior region of the vertebra. As can be seen in the images, there is a space beneath the endplate of a few millimeters which is not readily visible in two-dimensional images.



Figure 4.11 Demonstration of the extent of endplate fracture

# 4.4 Summary of Results

None of the specimens exposed to the smooth freeway cycling protocol demonstrated any visible damage to the intervertebral disc as brought on by the induced vibrations. All 15 injections (5 for each of the 3 discs) resulted in the disc being able to maintain intradiscal pressure (injections were made using thumb pressure applied against maximal resistance). Additionally, the nucleus pulposus remained within the area outlined by the safranin-O stain and no visible deterioration of the annulus fibers was detected upon dissection of the discs. The specimens did not have an endplate fracture on either the superior or inferior endplates adjacent to the intervertebral disc upon dissection. These results are in line with the preliminary study, where the two specimens exposed to the freeway cycling protocol showed no visible signs of damage to the intervertebral disc or any signs of endplate fracture upon dissection.

All three of the specimens exposed to the rough freeway cycling protocol demonstrated at least some signs of damage induced by cycling. Endplate fractures were found on the inferior of the two endplates adjacent to the intervertebral disc for all three specimens. All three specimens were unable to maintain intradiscal pressure during or before the third of the alcian blue injections (fifth injection overall). Two of the three intervertebral discs also showed signs of deterioration of the inner annulus and migration of either/both alcian blue stain and nuclear material beyond the inner annulus. The disc that did not demonstrate any damage to the annulus fibrosis was from a relatively young donor and was therefore of very good health. It should also be noted that all of the specimens exposed to the expansion joint cycling protocol during preliminary testing also had an endplate fracture along the inferior-most of the endplates adjacent to the intervertebral disc except the lone specimen exposed to loads at 0.5 Hz. The results for all

6 of the specimens used during final testing are summarized in Table 4.2, found below. Of particular interest is the consistency seen in the results within the two cycling protocols.

Specimen	Segment	Exposure	Annulus Damage?	Endplate Fracture?	When innerdiscal pressure was lost
044	L2-L3	Freeway			Never
067	L4-L5	Freeway			Never
073	L2-L3	Freeway			Never
033	L4-L5	Exp. Joint	~	~	2nd alcian injection
067	L2-L3	Exp. Joint		~	3rd alcian injection
073	L4-L5	Exp. Joint	~	~	2nd alcian injection

Table 4.2Final testing results

### **CHAPTER 5: Discussion**

### 5.1 Revisiting Final Testing Aims

Three goals were established for the final series of tests in section 4.1. These aims are discussed below in the context of experiment results.

1. Determine whether either of the two vibration signatures was injurious to the spine. Based on the results of the final round of tests, evidence supports the idea that the expansion joint exposure is more detrimental compared to the freeway exposure. Beyond this comparison, it is difficult to relate the exposures induced during cycling with what are seen in occupational settings because of the ex vivo nature of this study. The experimental model used was a body-disc-body model that was specifically chosen to reduce the number of variables and accentuate damage to the disc under cyclical loading. However, elimination of the posterior structures of the vertebrae (facet joints) as well as ligaments and muscles may have altered the effects of the vibrations. Additionally, the experiment relied on accelerating exposures into condensed periods of time – according to a spokesperson for King County Metro, a driver is not likely to drive on a freeway for more than 40 minutes at a given time and would encounter no more than 40-50 expansion joints on such a route. This very compressed duty cycle of the exposures also did not allow for any relaxation or healing the spine may undergo in vivo between exposures. Nonetheless, what can be said with confidence is, given the same conditions, evidence exists that the expansion joint exposure is more likely to induce damage to the human spine than the freeway exposure. A more definitive conclusion could be made by increasing the number of samples studied.

- 2. Determine whether the same trends of disc and lumbar body degeneration observed in the preliminary study occurred under the final protocol loading conditions. For the most part, the results of the dissection pointed to two distinct trends. Firstly, the specimens exposed to the freeway exposure showed no definitive signs of deterioration. Secondly, those FSUs exposed to the expansion joint segment developed endplate fractures on the inferior endplate adjacent to the intervertebral disc in addition to showing signs of damage to the annulus of the intervertebral disc. These results are consistent with what was seen in the preliminary tests.
- 3. *Outline the progression of disc damage.* Much of the deterioration observed within the annulus of the disc was found to occur in a non-specific direction with respect to the location of the nucleus. However, progression of injury could be detected to occur toward the exterior of the disc from the nucleus pulposus. This agrees with expectations based on the assumption (made during development of the staining technique) that damage would begin in the interior region of the disc and progress towards the exterior. The alternative means of injury progression (exterior of the intervertebral disc towards the interior) was not explored with this study protocol and is addressed in the study limitations section presented below.

#### 5.2 Study Limitations

The most obvious limitation of this study is the small sample size used. This is mostly due to the relatively limited availability of human specimens for laboratory testing and the necessarily strict requirement that the intervertebral disc be able to maintain intradiscal pressure. Unfortunately, the latter requirement eliminated half of the planned test specimens. Ideally, the same experiment could be performed with an increased sample size with paired specimens from the same donor. Putting each of the specimens from the same donor under differing exposures (as was done for two pairs of the final specimens) would strengthen this study. Additionally, if the specimens are not from the same level of the spine, similarly alternating FSUs from differing regions of the spine would also increase the validity of results since disc injuries occur throughout the lumbar region. As it stands, the rather small sample size prevents more definitive conclusions from being drawn with respect to study results.

An additional limitation with using human specimens is the age of the specimens. As can be seen from the results from specimen ID# 067, age can have a great impact on the health and overall quality of the disc. Because the supply of donor specimens is most always from older individuals, some preexisting deterioration of the intervertebral disc is unavoidable. Not only this, but bone quality could greatly impact whether or not an endplate fracture is encountered during testing. Specimens from elderly donors are more likely to have a decreased bone quality and may thus be more susceptible to the endplate fractures seen during testing.

A possible solution to the limitations surrounding the use of human tissues is to use an animal model. Though this could very well work for the validation of the staining techniques and the detection of damage to the hard bones of the vertebral bodies, the translation of displacement data collected on humans to animals, be they porcine or primate tissues, could pose issues related to the morphology of the spine of the groups studied. In particular, because quadrupedal animals do not axially load their spines, may have intervertebral discs which differ from humans in both shape and health. Additionally, bone strength may differ between the species as a difference in animal sizes would likely result in a difference in vertebral body size and specimens would

therefore have to be subjected to a proportionately different loading displacement for a fair comparison.

Another limitation of this particular study is that it only examines disc injuries which progress from the interior of the intervertebral disc towards the exterior. As previously stated, assuming disc injury progressed in one direction or the reverse was necessary for developing a practical staining protocol. The reverse means of disc injury progression (from the exterior of the disc towards the interior) was not tested in this study but could potentially be explored by immersing the FSU in a reservoir of stain during testing and then dissecting to determine if any stain was found to have penetrated the exterior of the annulus fibrosis. Such a test would not only involve large volumes of stain, but also require care to ensure that the equipment used in testing is remains undamaged from the cycling protocol. Also, it would be limited only to examining damage/degradation of the outer annulus. Endplate fractures induced by the cycling protocol (or that were preexisting) would be impossible to ascertain with this alternative staining technique.

The initial angle at which the FSUs are potted in the PMMA mold and subsequently loaded presents another limitation. Spine curvature and the angle at which individual vertebrae are at during rest vary greatly not only by region of the spine, but also between sitting and standing <sup>73-76</sup>. For this study, the specimens were potted at near-zero angles relative to the normal of the movement of the MTS actuator which mimics the angle of the intervertebral discs at the thoracolumbar junction when sitting. During sitting, however, lower lumbar segments tend to flex and bend relative to one another, thereby creating a nonparallel, angled alignment between adjacent vertebral bodies <sup>75, 76</sup>. Positioning FSUs at non-neutral angles greatly decreases the ability of the disc-body-disc segment to withstand applied forces without failure during loading cycles while better representing the position of the lower lumbar region of the spine when in a

seated position <sup>17, 77</sup>.

A final limitation of this study relates to the fact the displacement values used were derived from occupational exposures and not directly collected therefrom. For this project, the occupational displacements were collected in a vehicle, played into a hydraulic shaker, and then derived using, in part, a mathematical model. Additionally, the data collected also looked at the vibration across a very large region of the spine, rather than a single intervertebral disc. Using a more direct way of obtaining the occupational displacement of individual vertebral discs would be of interest in improving the accuracy of displacements of intervertebral discs as used within this study.

# 5.3 Future Directions

A potential improvement to the study would be to increase the number of road conditions analyzed. As this was a first-pass at developing techniques for conducting future studies of this sort, only two, very different road conditions were studied. Looking at roads traversed at lower speeds and with differing impulse signatures such as city streets and streets with pot holes or speed bumps may be of great future interest to better understand the development of LBDs across a range of WBV exposure signatures.

Another improvement that could be made to a follow-on study is the manner in which the intervertebral discs are sliced during dissection. Using a scalpel by hand tends to result in uneven cuts of the intervertebral disc and, in some cases, smearing of dye. Using a sharper tool with a longer blade to allow for single cuts would be of great help. Although efforts were made to identify such a tool, none were found that have been previously used in intervertebral disc discertions. A microtome could potentially solve this issue but because the discs contain liquid

stains, any freezing that may be required for proper slicing with the microtome may alter results. Additionally, pooling of stains may be liable to damage the microtome equipment.

Another potential improvement relates to the degree to which the results can be quantified. In this study, most results were based on qualitative visual inspection of the discs and dyes. Quantifying these observations for non-visual comparison across specimens is a desired but missing feature. Alternate studies following dye migration have relied on using digital image analysis computer programs to detail and compare images of intervertebral discs based pixels of stain in the annulus and of stain in the nucleus pulposus <sup>59</sup>. Such a method, however, is dependent upon the slice plane used during imaging and the image quality. As has been shown in results from this study, the concentration of stain tends to vary greatly at different levels of the intervertebral disc. As such, it seems an alternative method that takes this into account in some way must be developed.

A final direction towards which to take this project relates to comparing the vibration signatures applied working limits and/or industry standards. As previously stated, ISO standards exist regarding the limits of WBV exposure in a workplace. Determining how the induced cycling protocols compare to these WBV exposure limits would be of interest. Because these standards and limits often rely on the acceleration data from WBV sites, displacement waveforms could be doubly differentiated to obtain accelerations. Alternatively, accelerometers could be affixed to the actuator of the MTS as it oscillates for each of the two exposure signatures without applying a load to a FSU. This accelerometer data could then be processed and analyzed for comparison with working limits and standards.

#### 5.4 Conclusions

This thesis introduces methods which can be used as a basis for conducting more complete investigations into the effects of WBV on the development of LBDs. Based on a review of literature, this project was the first known to translate vibrations collected in an occupational setting into a laboratory setting to test on human spine specimens. Moreover, the study also introduced a novel staining protocol which employing readily available histological stains to outline the preexisting condition of the intervertebral disc and trace any induced by cycling the disc. Previous studies relied on radiography to establish the preexisting state of the intervertebral disc. Based on these two methodological innovations alone, this study stands to help advance knowledge regarding the etiology of LBDs as a result of WBV.

Based on the presented results, it can be said that evidence now exists that for an equal driving-time exposure, higher impulses from features of freeways such as expansion joints are much more likely to induce injury than the vibration induced from driving on a more even (smooth) driving surface. These injuries can be manifested as both endplate fractures as well as deterioration of the annulus of the intervertebral disc, which can lead to disc prolapse and/or herniation. However, the low sample number (n = 6) prevents more conclusive determinations from being made based on the results of this study.

Additionally, the presented results also provide evidence that progression of disc-related injury can proceed from the interior of the disc towards the exterior. Testing the alternative ("out-to-in") hypothesis could be done, as discussed in the study limitations. It should also be stated that these two modes of progression are not mutually exclusive – disc injury progression could very well progress simultaneously in both directions. For this experiment, increasing the number of samples would again allow for more definitive conclusions to be drawn.

Overall, this study demonstrates merit in terms of its methods and points towards conclusions that can be drawn with more extensive results. It can, in many ways, be considered a strong first step towards translating occupational exposures into a laboratory setting while also providing results that hint at the etiology and development of LBDs as a result of WBV.

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