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Mechanobiology of the Intervertebral Disc: Influences of Static and Dynamic Compressions on Energy Metabolism

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UNIVERSITY OF MIAMI

MECHANOBIOLOGY OF THE INTERVERTEBRAL DISC: INFLUENCES OF STATIC AND DYNAMIC COMPRESSIONS ON ENERGY METABOLISM

By

Chong Wang

A DISSERTATION

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MECHANOBIOLOGY OF THE INTERVERTEBRAL DISC: INFLUENCES OF
STATIC AND DYNAMIC COMPRESSIONS ON ENERGY METABOLISM

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Low back pain (LBP) is the leading reason for people to miss work and chief complaint of 5% people who visit doctors. Although it has a high impact in society, the causes of LBP cannot always be clarified. However, intervertebral disc (IVD) degeneration has been reported to be a possible leading source for LBP. Since IVD is the biggest avascular tissue in human body, nutrition supply plays very important role for IVD degeneration. IVDs are under mechanical load all the time in vivo, it is also been revealed by in vitro studies that mechanical loadings can alter cellular activities including metabolism. However, how mechanical loadings affect IVD metabolism has not been elucidated yet. Therefore, the purpose of this dissertation is to study the energy metabolism of IVD under compressive loadings.

A bioreactor system was built to apply static and dynamic mechanical loadings to IVD. A pump was utilized to maintain culture medium circulation during tissue culture and loading experiments; a load cell and a data acquisition device were also included in the system to record loading forces.

Adenosine-5’-triphosphate (ATP) is the “energy currency” of cell activities and extracellular ATP is an important molecule which can mediate various physiological activities by activating series of receptors. Therefore it is essential to detect
extracellular ATP contents. However, existing techniques have their drawbacks and are not ideal for in vivo/in situ measurement. We thus developed an optical biosensor for in situ measurement of extracellular ATP measurement in biological tissues. This ATP sensor utilized the technique of sol-gel coating and two layers of coatings were applied to the end of optical fibers; the first layer contained Ruthenium complex which can be excited by blue light with 465 nm of wavelength and emit red light with wavelength at 610 nm; the second layer contains two enzymes: glycerol kinase and glycerol 3-phosphate which can oxidase ATP and consume oxygen. During measurement, ATP molecules diffuse into the second layer and are oxidized thus decrease local oxygen concentration; this decrease of oxygen is detected by the first layer and is recorded by NeoFox® software; the signal detected has a linear relationship with the ATP content. This ATP sensor has a broad range of measurement at $10^{-3}$ mM to 1.5 mM. A compensation method was established to enable the measurement of ATP contents at different environmental oxygen concentrations. We also demonstrated that the performance of this sensor was not affected by environmental pH and derivatives of ATP such as adenosine diphosphate (ADP) and adenosine monophosphate (AMP) or adenosine.

From static and dynamic compression experiments of porcine IVDs it was found that under both static and dynamic compressive compressions, pH decreased and the contents of lactate and ATP increased significantly in both annulus fibrosus (AF) and nucleus pulposus (NP) regions, suggesting that compressive loading can promote ATP production via glycolysis and reduce pH by increasing lactate accumulation. We also detected high level of extracellular ATP contents in the NP region and compressive loadings significantly decrease extracellular ATP contents. Since ATP
can be utilized as intracellular energy currency and regulate a variety of extracellular activities through the purinergic signaling pathway, the findings of this dissertation suggest that compression mediated ATP metabolism could be a novel mechanobiological pathway for regulating IVD metabolism.
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This is for those I love and those who love me.
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CHAPTER 1: INTRODUCTION AND SPECIFIC AIMS

1.1 INTRODUCTION

There are many reasons for people to miss work. One of the leading contributors is low back pain (LBP), which is believed to affect 80% of the population at some point during their lifetime (Borenstein, Calin et al. 2012). Suffering from back pain is the chief complaint of 5% of people who visit the doctor in the US (2008). The total medical cost related to low back pain in the US exceeds 100 billion dollars every year (Willis 2009).

There are many factors that may cause LBP; such as spine stenosis, spinal lesions, and disc herniation. Doctors diagnose LBP through patients’ histories, imaging studies and physical tests. The equipment often employed in modern diagnosis of LBP includes discography, magnetic resonance imaging (MRI), computerized tomography (CT), x-ray imaging, bone scintigraphy (bone scanning) and electro-diagnostic assessments (such as needle electromyography, nerve conduction, electromyography (EMG) and evoked potential (EP) studies) as auxiliary methods of detecting LBP. The diagnosis and treatment technologies have been extensively developed over time. However, the cause of LBP is still unclear (Deyo 1998). Other non-disc sources such as sacroiliac joints or hip joints may also mimic back pain. Doctors must also be aware of other diseases such as Parkinson’s disease or Vitamin D deficiency (rickets) that can also mimic back pain.

While the exact cause of LBP remains unclear, recent studies revealed that intervertebral disc (IVD) degeneration is closely related to LBP (Luoma, Riihimaki et al. 2000; Roberts 2003). Although aging, nutrition supply, mechanical factors and
many other aspects contribute to disc degeneration, the mechanism of degeneration is not yet determined (Hadjipavlou, Tzermiadianos et al. 2008; Kandel, Roberts et al. 2008; Adams, Dolan et al. 2009).

IVD is the largest avascular tissue in human body and undergoes repeated static and dynamic loading. The structure and local mechanical environment of IVDs create energy pathways that rely on diffusion of small solutes from peripheral and vertebral blood vessels and convection (Urban, Holm et al. 1982) which are affected by many factors such as endplate calcification (Grunhagen, Shirazi-Adl et al. 2011). Nutrition supply is essential for cell activity and maintaining tissue integrity; therefore, it is important for IVD degeneration studies and regenerative methods development to elucidate the energy metabolism of IVDs under mechanical loading.

1.2 SPECIFIC AIMS

The following are the hypotheses of this study:

1. Mechanical loading affects energy metabolism of IVD;

2. Annulus fibrosus (AF) and nucleus pulposus (NP) tissues exhibit different energy metabolism.

To test these hypotheses, the following are the specific aims of this project.

Aim 1: To design and build a novel bioreactor system for whole IVD disc culture in order to perform static and dynamic compression studies.

Custom designed chambers will be fabricated, with a dialysis pump for disc culture. A bioreactor system will be built which is able to perform both static and dynamic (sinusoidal) compression experiments. Omega® load cell will be used to monitor load. Different frequencies of dynamic compression will be performed.
Aim 2: To develop novel optical sensors to detect glucose and ATP contents.

Blue LED light source, spectrometer and oxygen partial pressure monitoring software from Ocean Optics® will be used. Novel coating layers containing specific enzymes will be developed for detecting glucose and ATP contents.

Aim 3: To investigate the effects of static and dynamic compressions on energy metabolism of IVD.

In the static compression experiment, 10% strain will be applied to the IVD for 1 hour and 2 hours. In the dynamic compression experiment, IVD will be subjected to sinusoidal compressive strain of 15% at 0.1 and 1 Hz. After compression experiments, pH and the contents of ATP, glucose, lactate, and nitric oxide in AF and NP regions will be determined.

1.3 CONTENT OF THIS DISSERTATION

The overall objective of this dissertation is to investigate the energy metabolism of IVD under static and dynamic compressions, in order to better understand the mechanism of disc degeneration and its relation with low back pain. To achieve this goal, a bioreactor system with a step motor and a moving piece to perform static and dynamic compressive loads which was attached with a culture medium circulation system and was attached to a loading recording system was built to perform compressive mechanical loadings, both static and dynamic. A fiber optic based ATP sensor was fabricated using sol-gel entrapped enzyme technique for extracellular ATP measurement. Static and dynamic compression experiments were performed on porcine IVDs and the effects of compression on energy metabolism of two anatomical regions in the IVD were analyzed.

In Chapter 3, a custom-made bioreactor system is described. This system can perform controllable static and dynamic compressive loadings. A culture medium
circulation system consisting of a pump and a reservoir was attached to this bioreactor system for maintaining viability of IVD during experiments by circulating culture medium. A data acquisition device and a load cell were used to read and record the force response of IVD during compression experiments.

In Chapter 4, a novel optical fiber based ATP sensor is presented. This sensor utilizes the sol-gel entrapped enzyme technique, detecting ATP contents in target solutions or tissues by measuring local oxygen concentration change caused by enzymatic oxidization of ATP.

In Chapter 5, the changes of pH, lactate accumulation, total ATP content and extracellular ATP concentration in the IVD caused by static and dynamic loadings are illustrated. The difference of above measurements between AF and NP regions are also presented.

Chapter 6 is the peroration of this dissertation and recommendation of future works.

1.4 SIGNIFICANCE OF THIS STUDY

Low back pain in the US is the leading cause of lost working ability. Low back pain and related ailments cost more than $100 billion in the US every year (2008; Borenstein, Calin et al. 2012). Although the real causes remain to be elucidated, low back pain is closely associated with disc degeneration.

Due to the avascular nature of the IVD, nutrients are delivered via diffusion and convection over a long distance through the dense extracellular matrix to the cells. Poor nutrient supply has been considered a potential mechanism for disc degeneration (Urban and Roberts 2003; Urban, Smith et al. 2004; Stephan, Johnson et al. 2011). IVD cells consume glucose and oxygen to produce energy (i.e., ATP) which is essential for ECM biosynthesis pivotal for maintaining ECM integrity and preventing
disc degeneration (Hay 1991; Silva and Mooney 2004). Due to poor nutrient supply, ATP production may be a limiting factor for ECM biosynthesis in the IVD.

Since the main function of IVD is to support loads and to provide flexibility for the spine, IVDs are subjected to mechanical loadings frequently. Many mechanical events at high magnitude can be generated inside the IVD under mechanical loading, such as fluid flow, hydrostatic pressure, tension, deformation, shear force and streaming potential, etc. All of these have been proved to have effects on ECM synthesis of IVD cells (Ohshima, Urban et al. 1995; Ishihara, McNally et al. 1996; Handa, Ishihara et al. 1997; Hutton, Elmer et al. 1999; Iatridis, Mente et al. 1999; Lotz and Chin 2000; Chen, Yan et al. 2004; Maclean, Lee et al. 2004; Walsh and Lotz 2004). This indicates that ATP production may be affected by mechanical loading. In addition, cells constantly release ATP which can regulate a variety of cellular activities via purinergic receptors (Burnstock 2006). ATP released by IVD cells can be promoted by mechanical loading, including IVD cells (Fernando, Czamanski et al. 2011). Due to the avascular nature of IVD, ATP may accumulate extracellularly and regulate metabolic activities of IVD cells, including ECM production (Chowdhury and Knight 2006). Therefore, mechanical loading could influence biological function of the IVD by regulating intra- and extra- cellular ATP metabolism. The overall objective of the studies in this dissertation was to investigate the energy metabolism of IVD under static and dynamic compressions. Thus, the outcomes of the studies in this dissertation improve our understanding of cellular energy metabolism and nutrient transport in the IVD under mechanical loading and provide guidance for future studies which will investigate a new mechanobiological pathway to disc degeneration by governing ATP metabolism and develop new strategies to prevent or retard the degenerative process of IVD.
CHAPTER 2: BACKGROUND

2.1 STRUCTURE OF IVD TISSUE

IVD is a strong yet deformable tissue that lies between vertebrae and is important for the whole spine structure. The anatomic views of IVD are shown in Figure 2-1. IVD transfers loads and allows the spine to move through torsion, bending or compression (Hickey and Hukins 1980; Hadjipavlou, Tzermiadianos et al. 2008; Adams, Dolan et al. 2009). IVD is also one of the largest avascular cartilaginous tissues in the human body. The main methods for delivering nutrients into IVD are diffusion and convection.

There are three major anatomical parts of the IVD: the inner nucleus pulposus (NP), the outer annulus fibrosus (AF) and cartilaginous endplate (CEP) (Figure 2-1) (Kandel, Roberts et al. 2008). The composition of IVDs is not identical but differs according to different ages and locations within the spine (Pearce, Grimmer et al. 1987; Urban, Fairbank et al. 2001). However, in a healthy and normal disc, the NP is a jelly-like part which contains randomly oriented collagen fibrils suspended in a proteoglycan gel (Marchand and Ahmed 1990). The main content of NP is water; collagen (mainly type II and type VI) is randomly oriented to form a loose network and to hold proteoglycan (Wu, Eyre et al. 1987; Roberts, Ayad et al. 1991). Negatively charged glycosaminoglycan provides NP the osmotic properties, which enables NP to play an important role for distributing hydraulic pressure when mechanical loadings applied to the spine (Setton and Chen 2006).
Figure 2-1: Anatomy of the Intervertebral Disc
The AF is made of 15-25 distinct layers of organized collagen fiber lamellae which form the surrounding ring around the NP (Marchand and Ahmed 1990; N. 2005). Each lamella has collagen fibers (type II) aligning the same direction but has an angle of ~30° to adjacent lamellae. Lamellae have relatively loose connections to each other and this helps the IVD to transmit movement such as flexion and torsion (Bibby, Jones et al. 2001). The AF and endplate together confine the more flexible and deformable NP (Adam and Deyl 1984; Guiot and Fessler 2000).

The vertebrae body and IVD are separated by CEP, a thin layer of hyaline cartilage (Lundon and Bolton 2001). The CEP prevents the NP from being squeezed out under compression and allows loading to be evenly transferred to vertebral bodies (Ferguson and Steffen 2003). The CEP is porous to allow solute transportation through the endplate (Grunhagen, Shirazi-Adl et al. 2011); therefore, the CEP is a very important nutrition pathway for IVD. The AF can receive nutrition supply by solute diffusion and convection from surrounding blood vessels, while the CEP is the main route of nutrition supply for the NP. However, the nutrition pathway tends to be less effective as CEP calcification due to aging occurs (Figure 2-2).

Water makes up the largest portion of the IVD. Water makes up 60-70% weight of the AF and 70-90% weight for NP (Panagiotacopulos, Pope et al. 1987; Iatridis, MacLean et al. 2007). However, the water content varies due to age and region. Proteoglycan makes up 10-60% of dry weight of IVD tissue, collagens (all types) make up 15-65% dry weight and other matrix proteins make up 15-45% of dry weight in human IVDs (Hendry 1958; Kraemer, Kolditz et al. 1985; Panagiotacopulos, Pope et al. 1987; Eyre 1989; Johnstone, Urban et al. 1992; Pearce 1993; Gu, Mao et al. 1999). The composition of human IVD is summarized in Table 2-1.
Figure 2-2: Scanning electron microscopy of cartilage endplate from A) 30 years old and B) 80 years old human beings (Benneker, Heini et al. 2005).
2.2 CELL TYPES IN DIFFERENT IVD REGIONS

In addition to morphological differences in the AF and NP, they also have different cell types. Within the AF there are different types of cells from the outer region to the inner region and in the transition zone (TZ). The AF outer region cells are mainly fibroblastic and in each lamellae they orient along the predominant direction (Figure 2-3) (O'Halloran and Pandit 2007). The cells in the AF inner region and TZ have both chondrocyte-like and fibroblastic characteristics (Walker and Anderson 2004).

NP cells are larger than AF and TZ cells (Figure 2-4 & Figure 2-5); they are spherical and surrounded by capsules (Maldonado and Oegema 1992; Oegema 1993). The cells in the NP change over an individual’s lifetime. The bigger cells, which are predominant in the earlier ages, are notochordal. For human beings, mature NP cells begin to replace notochordal NP cells from the age of 10. The mature fibroblastic cells are round in shape and much smaller (Adams, Dolan et al. 2009; Guehring, Wilde et al. 2009). NP tissues containing notochordal cells usually maintain a water content between 80-88% while mature NP tissue is around 70% water (Humzah and Soames 1988). It has been suggested that the disappearance of notochordal cells is due to the insufficient nutrient supply and the change of NP cell type contributes to disc degeneration (Aguiar, Johnson et al. 1999).

The cell density in the AF is 7000 to 12,000 cells/ mm$^3$ and higher than NP with 3,300 – 4,400 cells/ mm$^3$ (Maroudas 1975). Cells only form 1% of the disc volume, however, they play pivotal roles in maintaining disc integrity and health by producing ECM components and the chemicals responsible for breaking down the matrix (Bibby, Jones et al. 2001).
Table 2-1: Human IVD composition summarized from literature (Hendry 1958; Kraemer, Kolditz et al. 1985; Panagiotacopulos, Pope et al. 1987; Eyre 1989; Johnstone, Urban et al. 1992; Pearce 1993; Gu, Mao et al. 1999). Other matrix proteins are also found in IVDs in small amount and thus not listed here.

<table>
<thead>
<tr>
<th></th>
<th>Annulus Fibrosus (AF)</th>
<th>Nucleus Pulposus (NP)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Cell density</strong></td>
<td>~9000 cells/mm$^3$</td>
<td>~4000 cells/mm$^3$</td>
</tr>
<tr>
<td><strong>Water (% wet wt.)</strong></td>
<td>60-70%</td>
<td>70-90%</td>
</tr>
<tr>
<td><strong>Proteoglycan (% dry wt.)</strong></td>
<td>10-20%</td>
<td>~50%</td>
</tr>
<tr>
<td><strong>Collagen (% dry wt.)</strong></td>
<td>50-70%</td>
<td>15-25%</td>
</tr>
</tbody>
</table>
**Figure 2-3:** Cross-section view of a porcine lumbar disc. Three distinct regions can be seen from the Figure: Annulus Fibrosis (AF), Nucleus Pulposus (NP) and Transition Zone (TZ).
Figure 2-4: AF cells from 6-month old porcine lumbar disc.
Figure 2-5: 6-month old porcine lumbar disc NP cells
These processes are highly energy demanding. Cells consume large amount of adenosine triphosphate (ATP), which is the main cell energy source and generated via glycolysis and the consumption of glucose and oxygen. IVDs are under daily cyclic mechanical loading in vivo; it is reported that mechanical loading can alter the consumption of oxygen and glucose, change ATP release in chondrocytes and promote ECM synthesis in vitro (Lee, Wilkins et al. 2002; Huang, Hagar et al. 2004; Chowdhury and Knight 2006; Fernando, Czamanski et al. 2011; Salvatierra, Yuan et al. 2011). This may indicate the change of overall energy production of the IVD cells (Urban, Smith et al. 2004). Since IVD becomes avascular a few years after the fetal stage, the nutrients supply of IVD is mainly by diffusion, which is not a very efficient pathway and can be affected by many factors (Urban, Smith et al. 2004). When this pathway is affected, the consequence is insufficient nutrient supply which causes decreased energy production and is the main contributing factor in disc degeneration.

In addition to the differences in cell type, AF has higher collagen content and lower proteoglycan content compared to NP. Due to the differences in cell type and structure, AF and NP may have different metabolism pathways (Raj 2008; Bron, Helder et al. 2009). However, research on the energy metabolism of IVD and possible relations with disc degeneration is limited. Previous studies have shown the difference in energy metabolism between AF and NP in an agarose gel model (Fernando, Czamanski et al. 2011; Salvatierra, Yuan et al. 2011). However, the in vivo environment involves the cell-ECM interaction and mechanical loading which can change the transport properties of the disc such as permeability and diffusivity.

2.3 NUTRITION SUPPLY AND IVD DEGENERATION

As mentioned above, IVDs are the biggest avascular tissue in the human body; the main nutrient supply pathways: diffusion and convection of solutes from periphery
blood vessels penetrated into outer lamellae and CEP are affected by factors such as diffusivity and tissue permeability. Furthermore, the number of periphery blood vessels decrease dramatically while the volume of IVDs grow rapidly around the age of three to four for human beings (Urban and Roberts 1995). Upon skeletal maturation remodeling of CEP initiates and leads to mineralization and bone eventually replaces cartilage. Change to the morphology of the CEP decrease the permeability; thereby decreasing the overall nutrient supply to the NP (Moore 2006). It has been shown theoretically and clinically that CEP calcification will significantly decrease glucose and oxygen concentration in NP region and initiate disc degeneration (Peng 2001; Jackson, Huang et al. 2011). IVDs also endure repeated mechanical loading. This loading, taken in combination with CEP calcification creates a harsh environment for IVD cells, especially NP cells (Guiot and Fessler 2000; Grunhagen, Shirazi-Adl et al. 2011). Consequently, fibro-chondrocyte-like mature cells will replace notochord cells in NP which is believed to protect NP cells from degradation and apoptosis (Erwin, Islam et al. 2011). Hence necrotic and apoptotic cells increase from about two percent of at fetal population to more than fifty percent in the adult as a result of a decrease in the number of notochord cells (Gruber and Hanley 1998).

It is believed that disc degeneration in humans starts at the beginning of second decade of life, earlier than any other musculoskeletal tissues (Miller, Schmatz et al. 1988; Boos, Wallin et al. 1993), and the severity increases along with age (Nerlich, Schleicher et al. 1997). Disc degeneration within a population increases with age: 20% of teenagers showed signs of disc degeneration, by age 50 the percentage increases to 97%. Furthermore, 10% lumbar discs are severely degenerated at age 50, compared to 60% at the age of 60 (Miller, Schmatz et al. 1988).
With age, the AF lamellae become thicker and more fibrillated. Cracks grow between lamellae and gradually disrupt the structure of the AF, decreasing the AF’s load bearing ability. The NP progressively loses its water content and morphology with age. Microscopically, collagen content of the NP increases and leads to a blurring of the transition zone that lies between AF and NP. These changes together may lead to IVD fracture. The color of IVD also changes with age (Figure 2-6), from white or transparency to yellowish or brown fragments (Urban and Roberts 1995).

IVD gains its mechanical strength from two major macromolecular components: proteoglycans and collagens. The network formed by collagen (mainly type I and type II fibrils) making up around 70% and 20% of dry weight in AF and NP, respectively, provide the tensile strength for IVD. Proteoglycan has a hydrophilic property and thus helps IVD to maintain a relatively higher water content (around 70% in AF and 90% in NP for normal IVDs) to endure compressions. Unfortunately, IVD gradually loses its proteoglycan and collagen content with age and aggravation of disc degeneration. It is reported that the most significant biochemical change detected in degenerated discs is the decreased content of proteoglycan (Eyre 1977). Collagen content also changes with disc degeneration but not as significantly as proteoglycans (Urban and Roberts 2003).
Enzyme activity is also an indicator of biochemical change of IVD degeneration (Urban and Roberts 2003). IVD cells not only produce molecules to synthesize ECM, but also produce enzymes (such as MMPs and aggrecanases) to break down ECM (Urban and Roberts 2003). The balance between synthesis and break-down is well maintained in the healthy discs, but it is disrupted in degenerated IVD. In the degenerated IVD, matrix metalloproteinase (MMP) and aggrecanases show higher levels of activity than in healthy discs.

Figure 2-6 shows the conditions of healthy and degenerated IVDs: healthy disc with a more hydrated NP and white color while degenerated disc lose much of water content in the NP region and has a yellow color. Degenerated discs usually have decreased heights which can be detected by clinical imaging techniques such as MRI and CT scan (Figure 2-7).
Figure 2-6: Sagittal views of human IVDs. Left: healthy from 20-30 year-old disc. Right: degenerated from 50-60 year-old disc (Adams, Stefanakis et al. 2010).
Figure 2-7: Different stages of human lumbar region disc degeneration obtained with Magnetic Resonance Images (MRI) (A) Shows a healthy disc with distinct NP and AF regions; (B) A disc in early stage of degeneration with slightly decreased disc height and reduced signal density of NP region. * indicates AF bulging inward. (C) A severe degenerated disc showing largely decreased height, * indicates large cleft (Smith, Nerurkar et al. 2011).
2.4 ENERGY METABOLISM OF IVD

Adenosine tri-phosphate (ATP) is called the “molecular unit of currency” for organism cells (Knowles 1980). Each ATP molecule contains three phosphate groups and can be hydrolyzed into adenosine di-phosphate (ADP) or into adenosine mono-phosphate (AMP).

The hydrolysis of one mole of ATP to ADP will release 30.5 kJ of free energy and hydrolysis of one mole of ATP to AMP will release 45.6 kJ of free energy (Berg, Tymoczko et al. 2012). For eukaryotic cells, many sources such as amino acids, fatty acids, carbohydrates and glucose can all be substrates for ATP production. Glucose is the most common one among all of these sources to produce ATP. However, fatty acids can be the primary energy source when high energy is required. Heart muscle is an example which utilizes fatty acids to produce the high amount of ATP required to maintain normal functions. The cell energy production process starts firstly when the substrate is transported into the cytosol where the energy metabolism is conducted; ATP is produced from a series of complex enzyme catalyzed reactions (Figure 2-8).

Two processes contribute to ATP production: aerobic respiration with adequate oxygen supply and glycolysis when oxygen is absent in the environment. Through aerobic respiration thirty-six (36) ATP will be produced per molecule of substrate; while for glycolysis three (3) net ATP will be yield per molecule of glycogen and two (2) net ATP will be produced per molecule of glucose (Lodish 2013).
Figure 2-8: Schematic representation of ATP production in cells (Siegel 1999).
Figure 2-9: Simplified flow of glycolysis (Lodish 2013).
It is reported that some cells such as immature IVD cells and chondrocytes contain glycogen-glucose deposits (Trout, Buckwalter et al. 1982). When necessary glycogen can be broken down to glucose and can be used to produce ATP. The intermediate of glycolysis, pyruvate, can enter mitochondria to undergo respiration for the production of ATP when oxygen supply is abundant. When oxygen supply is limited, lactate will be produced and only two (2) ATP per molecule of glucose will be yield from the reaction (Berg, Tymoczko et al. 2012; Lodish 2013). The overview of glycolysis is shown in Figure 2-9; in which ten intermediates are involved in a sequence of ten steps of reactions. The overall reaction with D-glucose as the initial substrate is as following:

Equation 1: Glycolysis Equation

\[
D - \text{Glucose} + 2[NAD]^+ + 2[\text{ADP}] + 2P_i \rightarrow \\
2[\text{Pyruvate}] + 2[NADH] + 2H^+ + 2\text{ATP} + 2H_2O
\]

Pyruvate and other intermediates that have been transported into mitochondria will be converted to Acetyl-coenzyme A, which is the initiating compound for the following citric acid cycle (TCA cycle, also referred as Krebs cycle) as shown in Figure 2-8. It is known that the Krebs cycle can produce 2 ATP molecules per cycle and also yield intermediates such as NADH, which is an important compound for oxidative phosphorylation or called electron transport chain. Oxidative phosphorylation also takes place in mitochondria and the location is the inner membrane. Mitochondrial respiration can theoretically produce 36 ATP molecules per glucose molecule.
2.5 EFFECTS OF MECHANICAL LOADING ON IVD CELLS

The main function of IVD is to provide flexibility and support load of the spine. Throughout the walking cycle the IVD undergoes frequent mechanical loading. Numerous mechanical events can be produced inside IVD by mechanical loading, such as fluid flow, tensile/compressive stress, shear force, matrix deformation, stream potential, hydrostatic pressure, etc. All these mechanical events were reported to regulate ECM synthesis of IVD cells (Ohshima, Urban et al. 1995; Ishihara, McNally et al. 1996; Handa, Ishihara et al. 1997; Hutton, Elmer et al. 1999; Iatridis, Mente et al. 1999; Lotz and Chin 2000; Chen, Yan et al. 2004; Maclean, Lee et al. 2004; Walsh and Lotz 2004).

High magnitude of static compression (> 1 MPa) has been shown to be detrimental to ECM synthesis and cell viability (Iatridis, Mente et al. 1999; Lotz and Chin 2000). Moderate static compression (0.1-0.4 MPa) can promote collagen and proteoglycan synthesis in explants and increase ECM gene expression under *in vitro* three-dimensional IVD cells culture system (Ohshima, Urban et al. 1995; Chen, Yan et al. 2004). It is showed that hydrostatic pressures lower than 3.0 MPa can increase synthesis of collagen and proteoglycan of IVD cells at *in vitro* three-dimensional culture or tissue explants (Ishihara, McNally et al. 1996).

It is found that dynamic compression of various frequencies and amplitudes can cause different responses from IVD cells. Previous studies showed that low frequency dynamic compression (e.g. 0.01 Hz, 1 MPa) applied to rat IVDs up-regulated anabolic gene expression (aggrecan, type I and type II collagen) in NP cells (Maclean, Lee et al. 2004).
Previous studies also demonstrated that mechanical loading influences cellular energy metabolism (Lee, Wilkins et al. 2002; Fernando, Czamanski et al. 2011; Salvatierra, Yuan et al. 2011). Hydrostatic pressure can affect cellular membrane transport of solutes (Benz and Conti 1986; Hall 1999) and glycolysis by changing the rate of glucose to lactate conversion (Takagi, Ohara et al. 1995). Hydrostatic pressure at physiological level (5-15 MPa) has been shown to up-regulate ECM synthesis in vitro by promoting incorporation rates of ECM building blocks (Hall, Urban et al. 1991). A recent study also showed that static compression can also inhibit the rate of oxygen uptake and lactate production of chondrocytes (Lee, Wilkins et al. 2002). It is also found that static and dynamic compressions promote ATP production and release from both AF and NP cells of IVD (Fernando, Czamanski et al. 2011; Salvatierra, Yuan et al. 2011). In summary, since ECM biosynthesis is a high energy demanding process (Lee and Urban 1997), the change of cell energy production caused by mechanical loads can lead to changes of cell metabolism.

2.6 EXTRACELLULAR ATP

ATP plays very important roles in regulating cellular activities other than serving as an energy molecule when released into extracellular space. Extracellular ATP (eATP) regulates many physiological activities by activating purinergic P2 receptors (ligand-gated ion channel) (Cook, Vulchanova et al. 1997) which are widely distributed in the nervous system, muscle, bone, endothelia and epithelia (Burnstock 2000; North 2002). ATP acts as a transmitter of the senses of pain (Cook, Vulchanova et al. 1997), mechanical loading (Cockayne, Hamilton et al. 2000; Vlaskovska, Kasakov et al. 2001), and temperature (Souslova, Cesare et al. 2000) by activating P2X receptors in the nervous system. ATP also plays an essential role in diseases, such as ischemia (Bush, Keller et al. 2000) and Parkinson’s disease (Przedborski and
In addition, ATP can initiate and modulate calcification of cartilaginous tissues (Felix and Fleisch 1976; Hsu and Anderson 1996; Golub 2011). It was found that 1 mM of ATP can promote tissue calcification, while higher ATP concentrations (e.g., 4 mM) can inhibit calcification (Hsu, Camacho et al. 1999). Therefore, the sensing of extracellular ATP \textit{in-vivo} has recently attracted increasing interest (Schneider, Egan et al. 1999; Brown and Dale 2002; Seminario-Vidal, Lazarowski et al. 2009; Trautmann 2009). However, the lack of efficient direct measurement techniques limits future studies of the comprehensive role of ATP. In addition, mechanical loading can promote ATP release from IVD cells (Fernando, Czamanski et al. 2011; Salvatierra, Yuan et al. 2011). Due to the avascular nature of the IVD, eATP may accumulate and has an influence on the biological function of the IVD.
CHAPTER 3 BIOREACTOR

3.1 PRINCIPLES OF BIOREACTOR

Human IVDs undergo different types of mechanical loading continuously; therefore, a bioreactor system is required to simulate these loadings to fulfill our specific aims of this study. One aim is to better simulate mechanical loadings applied to IVDs caused by human body movement. To fulfill these requirements, the design principle of this bioreactor system should cover the following points:

1. It should be able to work correctly in incubators with temperature, oxygen and carbon dioxide controls;
2. It should be able to perform both static and dynamic loadings;
3. It should be able to be connected to a culture medium circulation system.

We have designed and built a bioreactor system as shown in Figure 3-1. Figure 3-2 shows the bioreactor placed in an incubator with controlled temperature, oxygen level and carbon dioxide level. In Figure 3-3 details of a bioreactor chamber is shown.

This bioreactor system is driven by a SmartMotor® (Moog Animatics, Santa Clara, CA, USA), and SmartMotor® Interface (SMI) software was used to control the motor through a customer written program. An OMEGA® LC 402 High Accuracy Low Profile load cell (OMEGA Engineering Inc., Stamford, CT, USA) was utilized to measure the compression loadings applied to the IVD. NI USB 6590 digital data acquisition instrument (National Instruments Corporation, Austin, TX, USA) was connected to measure load response during compression; an OMEGA® (OMEGA Engineering Inc., Stamford, CT, USA) DP25P-C panel display was connected to
display the force readings. During disc compression experiments, the custom made chamber was connected with a Bio-Rad (Hercules, CA, USA) model EP1 Econo pump to maintain culture medium circulation (Figure 3-1 and Figure 3-2).

3.2 VERIFYING THE BIOREACTOR SYSTEM

3.2.1  Swelling and preload

IVDs undergo substantial mechanical loads *in vivo* and the external loads affect internal swelling pressure. The relationship of external loads to internal swelling pressure and proteoglycan contents affects fluid content of the IVD (Urban and McMullin 1988). Swelling may happen when the IVD lose weight bearing functionality *in vitro* (Urban and Maroudas 1981). Swelling of the IVD may cause leaching of proteoglycan and other components of the IVD and cause other deleterious consequences (Urban and Maroudas 1981; Holguin, Muir et al. 2007). Therefore, for *in vitro* culture of IVD, a preload is often added to the disc to balance the swelling pressure and thus prevent the IVD from swelling (Costi, Stokes et al. 2008). The effect of preload on mechanical properties of IVDs has previously been studied (Panjabi, Krag et al. 1977).

However, our preliminary study showed that porcine IVDs exhibited a very small disc heights change during overnight culture (2.3%±0.9%, n=3). Thus no preload was applied to the IVD during overnight culture in this study.
Figure 3-1: Bioreactor system.
Figure 3-2: Bioreactor in incubator
Figure 3-3: Bioreactor chamber with lid closed (up) and open (down).
3.2.2 Verifying displacement-controlled dynamic loading

The dynamic compressive loading applied in this study was displacement-controlled. An initial 10% strain was added to the disc by 10 steps with 1% per step. Then a dynamic loading with 10% strain was applied. Therefore the maximum strain applied on the disc during the experiment was 15% (Figure 3-4). The range of strain during dynamic compression experiments was 5%-15%.

IVDs have high fluid content which will be squeezed out slowly when compressive loads are applied. Therefore the IVD has a delayed response to compressive loads applied which may cause a so-called “lift off” effect. This effect indicates the status: when the moving piece that applies compression to the IVD is moved upward, the IVD is not able to follow due to the delayed response. In this short period of time, the moving piece and the IVD will be out of contact, the disc will thus not experience the next compression cycle. To verify if this effect happens on our bioreactor, we used a load cell with an attached data acquisition device, a display device and a recording program on the computer to monitoring the loading force fluctuation during the compression. A typical 10-second period of loading force approaching the end of 1-hour dynamic compression was shown in Figure 3-5. During the dynamic compression experiments, the force detected by the load cell never fell under zero (0). Therefore we have not observed any lift off effect on our bioreactor.
Figure 3-4: Schematic diagram of displacement-controlled dynamic loading.
Figure 3-5: 10 seconds of loading force-time recorded for 1 Hz dynamic compression.
4.1 INTRODUCTORY REMARKS

Adenosine-5'-triphosphate (ATP) is an important multi-functional molecule which was first discovered in 1929 by Karl Lohmann and has long been recognized as “the molecular unit of currency” (Lohmann 1929; Knowles 1980). ATP has two phosphate groups which release energy when hydrolyzed, and this is the energy source for the metabolic activities of ubiquitous organisms (Knowles 1980). When it is released to the extracellular environment, ATP can mediate numerous physiological activities by activating purinergic P2 receptors (ligand-gated ion channel) (Cook, Vulchanova et al. 1997) which are widely distributed in the nervous system, muscle, bone, endothelia and epithelia (Burnstock 2000; North 2002). In the nervous system, ATP acts as a transmitter of the senses of pain (Cook, Vulchanova et al. 1997), temperature (Souslova, Cesare et al. 2000), and mechanical loading (Cockayne, Hamilton et al. 2000; Vlaskovska, Kasakov et al. 2001) by activating P2X receptors. ATP also plays an essential role in diseases, such as particular ischemia (Bush, Keller et al. 2000) and Parkinson’s disease (Przedborski and Vila 2001). In addition, ATP can initiate and modulate calcification of cartilaginous tissues (Felix and Fleisch 1976; Hsu and Anderson 1996; Golub 2011). It was found that 1 mM of ATP can promote tissue calcification, while higher ATP concentrations (e.g., 4 mM) can inhibit calcification (Hsu, Camacho et al. 1999). Therefore, the sensing of extracellular ATP in-vivo has recently attracted increasing interest (Schneider, Egan et al. 1999; Brown and Dale 2002; Seminario-Vidal, Lazarowski et al. 2009; Trautmann 2009). However, the lack
of efficient direct measurement techniques limits future studies of the comprehensive role of ATP.

The Luciferin-luciferase method for measuring ATP content is well known for its high sensitivity and has been widely applied for ATP analysis (Rong, Gourine et al. 2003). However, it is only feasible to apply this method for ATP measurement in solution (Brown and Dale 2002). Recently, amperometric ATP biosensors were developed to detect ATP using multi-enzymatic reactions (Katsu, Yang et al. 1994; Compagnone and Guilbault 1997; Kueng, Kranz et al. 2004; Llaudet, Hatz et al. 2005). The biosensor containing glucose oxidase and hexokinase was able to determine ATP concentration in the presence of glucose with a detection sensitivity of 10 nM (Kueng, Kranz et al. 2004). In that biosensor, glucose was catalyzed by hexokinase in the presence of ATP, which reduced the production of hydrogen dioxide from oxidation of glucose by glucose oxidase. Recent studies developed amperometric ATP biosensor based on sequential enzymatic reactions of glycerol kinase (GK) and glycerol 3-phosphate oxidase (G3POX), which broke down ATP and produced hydrogen dioxide (Katsu, Yang et al. 1994; Llaudet, Hatz et al. 2005). In those biosensors, ATP concentration was determined by measuring electrical current using a polarized platinum electrode which oxidized hydrogen dioxide to oxygen and water. However, since those amperometric biosensors can be affected by many other compounds that naturally exist in biological tissues such as ascorbate and glucose (Kueng, Kranz et al. 2004; Llaudet, Hatz et al. 2005), it could potentially be problematic if they are used for in-vivo measurements. To avoid this problem, in this study, an optical ATP biosensor was developed using an optical oxygen sensing technique, based on fluorescence quenching of excited ruthenium complexes, which determined changes in oxygen concentration during enzymatic ATP breakdown. A
compensation method was also developed to enable the new ATP optical biosensor to determine ATP concentration at different oxygen levels after calibration tests.

4.2 EXPERIMENTS

4.2.1 Materials

Tetramethoxysilane (TMOS), methyltrimethoxysilane (Me-TriMOS), GK, G3POX, poly (ethylene glycol) (PEG), glycerol and Tris (4, 7-diphenyl-1, 10-phenanthroline) - ruthenium (II) bis (hexafluorophosphate) complex were obtained from Sigma (St. Louis, MO, USA).

Adenosine 5’-triphosphate sodium salt purchased from Sigma (St. Louis, MO, USA) was used for making standard solution. Cyanoacrylate glue was obtained from Pacer Technology (Rancho Cucamonga, CA, USA). NeoFox® phase measurement systems were purchased from Ocean Optic Co. (Dunedin, FL, USA). L-ascorbic acid was purchased from Sigma (St. Louis, MO, USA).

4.2.2 Fabrication of ATP optical sensor

(a) Chemical mechanism

The mechanism of the ATP optical biosensor developed in this study was based on the following sequence of enzymatic reactions (Murphy and Galley 1994):

Equation 2: Enzymatic Reaction 1

\[ \text{ATP + Glycerol} \xrightarrow{\text{GK}} \text{Glycerol-3-phosphate} + \text{ADP} \]

Equation 3: Enzymatic Reaction 2

\[ \text{Glycerol-3-phosphate} + \text{O}_2 \xrightarrow{\text{G3POX}} \text{Dihydroxyacetone phosphate} + \text{H}_2\text{O}_2 \]
As shown in these enzymatic reactions, ATP can be catalyzed by GK to ADP in the presence of glycerol and yields glycerol-3-phosphate which is further oxidized into dihydroxyacetone phosphate by G3POX. During the second chemical reaction, oxygen is consumed. Therefore, the ATP concentration can be determined by measuring changes in oxygen concentration.

(b) Design of the ATP optical sensor

The ATP biosensor was fabricated by coating two sol-gel layers on the tip of an optical fiber (Figure 4-1). In the first layer (i.e., oxygen sensing layer), ruthenium complexes were entrapped in silicate matrix which was fixed to the end of the optical fiber using cyanoacrylate glue. This layer was used to detect change in oxygen concentration. On top of the first layer, a layer of silicate matrix containing G3POX and GK was coated (i.e., enzyme layer), yielding the ATP optical biosensor. Glycerol and PEG were also included in the silicate matrix to stabilize the activity of the enzymes (Llaudet, Hatz et al. 2005).

During the measurement, a semi-permeable silicate matrix allowed ATP to diffuse into the enzyme layer and induce enzymatic reactions described in the reactions Equation 2 and Equation 3. Due to chemical reactions, changes in the oxygen concentration in the silicate matrix can alter the quenching of ruthenium complexes and thus change the decay time of fluorescence emission (Lippitsch, Pusterhofer et al. 1988; Lakowicz 2006). A NeoFox® phase measurement system was used to determine the decay time of fluorescence emission of ruthenium complexes in this study. The precision of decay time measurement is 0.000084 μs. To test and calibrate the ATP sensor, a linear relationship between the inverse of decay time and the standard ATP concentration was examined (see the section on calibration test).
(c) Preparation of the oxygen sensing layer

The oxygen sensing layer was prepared based on a modified version of the protocols previously described in literature (McEvoy, McDonagh et al. 1996; McNamara, Li et al. 1998; Xiong, Xu et al. 2010). Briefly, a silicate solution was made by mixing 7μl of Me-TriMOS, 13 μl of TMOS, 60μl of de-ionized water and 1μl of 40 mM HCl. The ratio of the reagents is essential for coating pore size and morphology. Then the silicate solution was sonicated for at least 15 minutes and stirred for 1 hour. These processes are critical to yield a firm and crack-free coating layer. A solution of ruthenium complexes was made by dissolving ruthenium complexes in methanol (5.4 μg/μl). This ruthenium complex solution was mixed with equal amounts of the silicate solution and phosphate buffer solution (PBS) at pH 7.5.

Immediately after applying a very thin layer of cyanoacrylate glue on the end of the optical fiber, one microliter of the ruthenium complex-silicate mixture was spread on the top of the cyanoacrylate glue layer. Cyanoacrylate glue was used to provide a firm adhesion of the ruthenium complex-silicate layer on the optical fiber and prevent detachment of the coating during measurement. The coated optical fiber was left in ambient conditions for 10 minutes before being stored at 4°C overnight. Once the coating became solid, the probe was ready for coating the second enzyme layer or was used for measurement of oxygen concentration.

(d) Preparation of the enzyme layer

Similar to the oxygen sensing layer, a silicate solution was prepared as described in the previous section and used to entrap enzymes in the second layer (Figure 4-1). GK and G3POX were dissolved in PBS with a supplement of PEG and glycerol. The enzyme concentration depends on the ATP concentration range to be measured, varying from 500-1500 units/ml of each.
PEG is well known for its ability to form crosslinks (Kulkarni, Hukkeri et al. 2005) which help to immobilize enzymes and maintain enzymatic activities. The silicate solution and the enzyme solution were then mixed at a 1:1 ratio. One microliter of the enzyme-silicate mixture was immediately spread on top of the oxygen sensing layer to complete the ATP biosensor. The biosensor was ready for ATP measurement after overnight storage in a 4°C environment away from light. To maintain enzymatic activity, the biosensor was stored in PBS with 2mM glycerol after re-hydration.

4.2.3 Calibration test of the ATP sensor

Oxygen sensing is based on the fluorescence quenching of ruthenium complex in the presence of oxygen which can be described by the Stern-Volmer equation:

Equation 4: Stern-Volmer equation

\[
\frac{\tau_0}{\tau} - 1 = K_{sv}[O_2]
\]  

Where \(\tau_0\) and \(\tau\) are the decay times in the absence and presence of oxygen, respectively, \(K_{sv}\) is the Stern-Volmer quenching constant, and \([O_2]\) is the oxygen concentration. The decrease in the oxygen concentration \((\Delta [O_2])\) due to the enzymatic reactions was assumed to be linearly proportional to the ATP concentration ([ATP]):

Equation 5: Relationship between oxygen concentration and ATP concentration

\[
\Delta [O_2] = k[ATP]
\]  

Here \(k\) is the enzymatic reaction parameter which is a function of initial oxygen level and enzyme concentration.
Figure 4-1: Cross sectional view of the ATP sensor
In the presence of ATP, the quenching of ruthenium complex in the coating can be described by Equation 6: Ruthenium quenching in the presence of ATP

\[
\frac{\tau_0}{\tau} - 1 = K_{sv} ([O_2] - \Delta [O_2])
\]  

(3)

When the reference oxygen level ([O_2]) and the enzyme concentration in the coating remain constant, the enzymatic reaction parameter k can be assumed to be a constant. Based on Equation 6, a linear relationship between the inverse of decay time and the ATP concentration can be derived as:

Equation 7: Relationship between ATP concentration and decay time

\[
[ATP] = A \frac{1}{\tau} + B
\]  

(4)

Where \( A = -\frac{\tau_0}{K_{sv} k} \) and \( B = \frac{1}{K_{sv} k} + \frac{1}{k} [O_2] \).

In the calibration test, the ATP biosensor was connected to the NeoFox phase measurement system which determined the decay time of fluorescence emission at the wavelength of 600 nm by exciting ruthenium complexes with an impulse light (44 kHz) at the wavelength of 465 nm (Figure 4-2). The resolution of decay time measurement was 10-4 sec. The biosensor was immersed in the PBS containing 2mM glycerol in a beaker. A magnetic stirring bar was used to stir the solution to distribute all solutes uniformly. After a reference decay time was established, a stock ATP solution of 1mM was added to achieve designated ATP concentrations in the PBS. The decay time was measured for individual ATP concentrations to determine the linear relationship described in Equation 7. The stability of the ATP sensor was also studied. The responses of three sensors to zero and 200 μM ATP were measured for five consecutive days.
The sensors were stored in PBS contain 2 mM glycerol at 4°C between measurements. The stability of the ATP sensor was analyzed by comparing the difference in the inverse of decay time between the responses to zero and 200 μM ATP.

The stability of the ATP sensor was also studied. The responses of three sensors to zero and 200 μM ATP were measured for five consecutive days. The sensors were stored in PBS contain 2 mM glycerol at 4°C between measurements. The stability of the ATP sensor was analyzed by comparing the difference in the inverse of decay time between the responses to zero and 200 μM ATP.

4.2.4 Examination of the effects of pH on the ATP biosensor

pH in biological tissues may be different from that in the in-vitro calibration condition. This difference may affect the in-situ measurement of the biosensor. Therefore, it is necessary to either calibrate the biosensor under the same conditions as biological tissues or develop a method to compensate for the possible effect that might be introduced by the difference in pH. In this study, the effect of pH on the biosensor measurement was examined by adding an acidic solution to alter the pH in the PBS solution during the calibration test described in the previous section. Then, the measurements of the ATP biosensor at different pH was recorded and normalized for comparison.
**Figure 4-2**: Mechanism of optical biosensor system.
4.2.5 Examination of the effects of ADP, AMP, adenosine, and ascorbate on the ATP biosensor

Extracellular ATP is often hydrolyzed into ADP, AMP, or adenosine by ectonucleotidases (Meghji, Pearson et al. 1992). Those adenine nucleotides may coexist with ATP in biological tissues. In addition, previous studies showed that ascorbate affects ATP measurement of amperometric biosensors (Katsu, Yang et al. 1994; Kueng, Kranz et al. 2004; Llaudet, Hatz et al. 2005). Therefore, the effects of ADP, AMP, adenosine and ascorbate on the ATP biosensor were examined on three sensors in this study.

4.2.6 Compensation of the ATP biosensor under different oxygen levels

As described in Equation 7, a linear relationship between the inverse of decay time and the ATP concentration can be established at the same initial oxygen level. Due to differences in tissue material properties, tissue oxygenation varies among different biological tissues. In order to measure ATP concentration under different oxygen levels, a compensation method was developed based on the following mathematical background. It was assumed that the chemical reaction parameter $k$ in Equation 7 is a function of the oxygen level $[O_2]$:

Equation 8: Relationship between $k$ and oxygen level

$$k = C[O_2] + D$$

(5)

Here $C$ and $D$ are the material constants. After $K_{sv}$, $\tau_0$, $C$, and $D$ are experimentally determined, Equation 7 can be used to determine the ATP concentration (i.e., $[\text{ATP}]$) based on the measurements of the oxygen level (i.e., $[O_2]$) and the delay time (i.e., $\tau$). To test this compensation method, $K_{sv}$, $\tau_0$, $C$, and $D$ for the ATP biosensor were determined from two calibration tests conducted at the oxygen levels of 10% and 20%.
Experimental data obtained from another calibration test at the oxygen level of 15% were used to verify the compensation method. An oxygen regulated incubator was used to provide desired oxygen conditions.

4.2.7 *In situ measurement of extracellular ATP in porcine intervertebral disc*

To verify the ability of the ATP biosensor for measuring tissue ATP concentration, a tissue measurement experiment was conducted. Porcine NP tissue was obtained from 8-12 month old pig and NP tissue was extracted. This NP tissue was frozen for one day and then immersed into PBS overnight to make sure all ATP was released into the solution. Then this NP tissue was washed three times using PBS and transferred into another PBS solution containing 200 μM ATP. After one hour the tissue was taken out and placed in a petri dish, the ATP sensor was inserted to measure the ATP concentration. The measurement was compared with the real ATP concentration to verify the ability of the sensor for tissue measurement. We found that our sensor has the ability to accurately measure tissue ATP content: the measured ATP contents are 103%±4.1% to the real ATP content (n=3).

To test the capability of the ATP biosensor to measure ATP levels in biological tissue; the measurement of extracellular ATP was performed on lumbar intervertebral discs of 12 month-old pigs obtained within 3 hours of sacrifice. Functional spinal units (FSUs) were isolated as described in our previous study (Fernando, Czamanski et al. 2011). The harvested FSUs were then placed and cultured in custom built chambers overnight at 37°C with continuous circulation of culture medium. After overnight culture, a transversal cut was made at the mid-disc height of the FSUs and the nucleus pulposus region was exposed to the ambient condition (20% oxygen level). In order to avoid temperature effect, the ATP measurement was performed using the
ATP biosensor when the temperature of the tissues was reduced to the same level as the calibration test of the sensors (i.e., ambient temperature). The temperature of tissues was monitored using a thermometer. An extra sensor was prepared without the second enzyme layer for determining in-situ oxygen level at the same locations. Calibration of the sensors was performed at the oxygen levels of 10% and 20%. In addition, based on our pilot study, the decay time returned to original value at zero ATP level after the sensors were washed in PBS solution with stirring for 10 minutes between measurements. This was applied to prevent carry over effect.

4.3 RESULTS

4.3.1 Calibration test of the ATP biosensor

In the calibration test, when ATP diffused into the enzyme layer of the biosensor, catalysis of ATP decreased local oxygen concentration and thus increased the decay time of fluorescence emitted by the ruthenium complexes. Typical response and calibration curve of the ATP biosensor for different ATP concentrations are shown in Figure 4-3. The rising part of the decay time in the response curve of the ATP biosensor reflected diffusion of ATP into the enzyme layer and initiation of the chemical reactions which reduced the local oxygen partial pressure to a certain level (Figure 4-3a). The inverse of decay time was linearly proportional to the ATP concentration (Figure 4-3b) as described in Equation 7. As shown in Figure 4-3b, the calibration was performed 3 times within 6 hours and small standard deviations indicated the sensor had a good repeatability. By changing the concentration of enzymes, the ATP sensor can be used for different measuring ranges. Based on our experiments, the ATP biosensor had a low detecting limit of $10^{-3}$ mM (Figure 4-4a) and saturated at 1.5 mM (Figure 4-4b).
Figure 4-3: (a) A typical decay time curve of the ATP sensor in response to the ATP concentrations of 50, 100, 200, and 300 μM and (b) the corresponding calibration curve (mean values ±standard deviation).
Figure 4-4: (a) A typical sensor response at low ATP levels (1, 5, 10 and 20 μM) and (b) a representative calibration curve showing a linear relationship between the inverse of decay time and the ATP concentration up to 1.5 mM.
4.3.2 The effects of ADP, AMP, adenosine, ascorbate and pH on the ATP biosensor

To examine the effect of ADP, AMP, and adenosine, three ATP biosensors were first equilibrated in the PBS solution without ATP and other adenine nucleotides and then a stock solution of ADP, AMP, or adenosine was added to a final concentration of 200 mM in the PBS and the decay time was recorded. It was found that the ATP biosensors developed in this study exhibited no responses to adenosine, AMP and ADP (Figure 4-5). Similarly, ascorbate (400 μM) did not exhibit any effect on the ATP biosensors (data not shown).

For the pH test, the responses of three ATP biosensors were tested at the ATP concentration of 100 μM. The values of pH were chosen based on pH in the normal organism tissues which mostly varies from 6.5-7.5 (Gerweck and Seetharaman 1996). No significant effects of pH were found on the ATP biosensors (Figure 4-6). This is consistent with the previous study (Llaudet, Hatz et al. 2005).
4.3.3 Compensation of the ATP biosensor under different oxygen levels

A linear relationship between the parameter k and the oxygen concentration was found between 10% and 20% oxygen levels (Figure 4-7b) as described in Equation 8. Furthermore, the errors in the ATP measurement by the compensation method for the oxygen level of 15% were found to be less than 5% at different ATP levels (Figure 4-8).

4.3.4 Sensor stability

The responses of three sensors to 200 μM ATP solution were examined for five consecutive days. The signal strength of florescence, which is defined as the difference between the inverse of decay times measured at 0 and 200 μM ATP, gradually decreased and fell to about 65% at day 5 compared to day 1 (Figure 4-9). This may mainly attribute to the diffuse out of enzyme loosely entrapped at the top of the coating layer (Kueng, Kranz et al. 2004).
Figure 4-5: The typical reaction of the ATP sensor to ADP, AMP and adenosine of 200 μM compared to ATP of 100 μM.
Figure 4-6: Normalized signal of the ATP sensor in response to 100μM ATP at different pH conditions (n=3). The data were normalized to the response at pH 7.5 and are expressed as mean values ± standard deviation.
**Figure 4-7:** (a) Typical calibration curves obtained at different oxygen concentration and (b) the corresponding linear relationship between the parameter $k$ and oxygen concentration.
Figure 4-8: Accuracy of the oxygen compensation method on determination of ATP concentration at 15% oxygen level using the parameters obtained from the calibration tests at 10% and 20% oxygen levels (n=3). The data were normalized to the corresponding ATP concentration and expressed as mean values ± standard deviation.
Figure 4-9: Stability of the ATP biosensor. The signal strength of fluorescence is defined as the difference between the inverse of decay times measured at 0 and 200 μM ATP. The data are expressed as mean values ± standard deviation.
4.3.5 Measurement of extracellular ATP in porcine intervertebral disc

To verify the ability of the ATP biosensor for measuring tissue ATP concentration, a tissue measurement experiment was conducted. Porcine NP tissue was obtained from 8-12 month old pig and NP tissue was extracted. This NP tissue was frozen for one day and then immersed into PBS overnight to make sure all ATP was released into the solution. Then this NP tissue was washed three times using PBS and transferred into another PBS solution containing 200 μM ATP. After one hour the tissue was taken out and placed in a petri dish, the ATP sensor was inserted to measure the ATP concentration. The measurement was compared with the real ATP concentration to verify the ability of the sensor for tissue measurement. We found that our sensor has the ability to accurately measure tissue ATP content: the measured ATP contents are 103%±4.1% to the real ATP content (n=3).

Typical responses of the oxygen and ATP sensors during the extracellular ATP measurement in the nucleus pulposus region of the porcine intervertebral disc were shown in Figure 4-10. After the tissue was exposed to the ambient condition (i.e., 20% oxygen) for a certain time (about 10 min), the oxygen level was still less than the ambient oxygen level (20%) shown in Figure 4-10. The differences in oxygen measurement between the ATP and oxygen sensors indicated oxygen consumption by enzymatic ATP breakdown and were used to determine ATP concentration using Equation 5 with k determined from the calibration tests at 10% and 20% oxygen levels. The extracellular ATP concentration in the nucleus pulposus region of the porcine intervertebral disc was 190 ± 30 μM (n=3).
Figure 4-10: Typical responses of the oxygen and ATP sensors during the extracellular ATP measurement in the porcine intervertebral disc.
**4.4 DISCUSSION**

This study successfully developed a new optical biosensor for in-situ ATP measurement, which was demonstrated by the calibration test and the extracellular ATP measurement in the porcine intervertebral disc.

The optical ATP biosensor utilizes the principle of fluorescence emission of ruthenium complexes which does not exhibit any common electro active interferences by naturally occurring compounds (e.g., ascorbate) as described in previous studies of amperometric biosensors (Katsu, Yang et al. 1994; Kueng, Kranz et al. 2004; Llaudet, Hatz et al. 2005). Therefore, the optical ATP biosensor provides a good alternative method for ATP measurements especially for in-vivo and in-situ measurements.

When the optical ATP biosensor is immersed in a solution containing certain ATP levels, ATP diffuses into the coating layer through the microspores and is catalyzed by enzymes. The process of diffusion is driven by a concentration gradient. Since the concentration gradient is determined by normalizing the concentration difference between the solution and the enzyme layer of the ATP biosensor by the thickness of coating, the response time of the biosensor inversely depends on the coating thickness. Therefore, the response of the ATP biosensor can be further improved by decreasing the thickness of coating. Furthermore, since the rate of oxygen consumption in the enzymatic reactions depends on the concentration of enzymes, the sensitivity of the ATP biosensor can be enhanced by increasing the enzyme concentration in the coating layer.

In the extracellular ATP measurement of porcine intervertebral discs, the oxygen level in the tissue still remained low (Figure 6) after the temperature of the tissue was equilibrated with the ambient condition (i.e., 25°C and 20% oxygen level). It suggested that the oxygen solubility in biological tissues is different from the
condition in the calibration test. Therefore, the oxygen compensation method established in the study is a key component for precise in-situ ATP measurement. By using this method, the optical ATP biosensor was able to determine the extracellular ATP content in the intervertebral disc.

The ambient air temperature for calibration is often around 25 °C and the body temperature is around 37 °C. Since the principle of oxygen sensing using ruthenium complexes involves energy transfer, previous studies have demonstrated that Stern-Volmer quenching constant \( K_{sv} \) and \( \tau_0 \) are dependent on temperature (Lakowicz 2006; Morris, Roach et al. 2007). In addition, the enzyme activities could be affected by temperature. Therefore, the temperature effects on \( k \), \( \tau_0 \) and \( K_{sv} \) need to be compensated if the temperature at the measurement sites is different from that in the calibration test.

Luciferin-luciferase assay has long been used for determining the extracellular ATP content (Leach 1981). This method has many advantages such as ability to detect ATP at low concentration (Spielmann, Jacob-Müller et al. 1981). However, Luciferin-luciferase assay also requires strict conditions and operations and many factors can cause contamination to the assay which limits its application in in-vivo/in situ measurements. Additionally, Luciferin-luciferase assay is not feasible for real time monitoring ATP content change (Lundin 2000). Electrochemical ATP sensors with high sensitivity were developed by other researchers (Kueng, Kranz et al. 2004; Llaudet, Hatz et al. 2005). However, these sensors measured ATP by detecting current produced during chemical reaction and thus can be affected by other charged molecules such as ascorbic acid and urinate (Miele and Fillenz 1996). For example, ascorbate has a concentration range of 200-400 mM in human tissue and can generate up to 100 nA of current when using an electrode with 0.5 mm length and 50
μm diameter (Rice 2000; Llaudet, Hatz et al. 2005). In addition, the chemical reaction required in electrochemical ATP sensors may be affected by substances in body tissues. For instance, the ATP sensor developed by Kueng et al. requires glucose to be present at a known concentration. In human tissues, however, glucose content varies from different tissue and at different time points during a day (Maggs, Jacob et al. 1995; Daly, Vale et al. 1998). These factors make electrochemical ATP sensors difficult to be used for in vivo/in situ ATP measurement. The design of our sensor, however, has advantages in in vivo/in situ ATP measurement and potential for real time monitoring changes in ATP level. Furthermore, it was reported that that extracellular ATP concentration can reach several hundred micro molar at tumor interstitium (Pellegatti, Raffaghello et al. 2008), while extracellular ATP concentrations are around 4 mM in rabbit central nervous system during systematic inflammatory response (Gourine, Dale et al. 2007). Therefore, the new ATP sensor developed in our study can be used to monitor ATP level in those in-vivo conditions.

A new optical ATP biosensor was successfully developed with less chemical interference. The compensation method was also established to enable the new biosensor to detect ATP at different oxygen levels. This study demonstrated that the newly developed optical biosensor is feasible for in-situ extracellular ATP measurement.

4.5 OTHER OPTICAL SENSORS

Other optical sensors including oxygen sensor and glucose sensor were fabricated using similar mechanism with ATP sensor described above. Only the coating layer containing Ruthenium complex is applied to optical fiber to fabricate oxygen sensor; the coating layer is modified to be able to sustain under direct contact with tissues. G3POX and GK was replaced by glucose oxidase (Sigma Aldrich, St. Louis, MO,
USA) to yield glucose sensors as described by previous literatures (Wolfbeis, Oehme et al. 2000; Endo, Yonemori et al. 2006). These sensors were also used to measure oxygen content and glucose content distribution in IVD as shown in Chapter 5. The enzyme catalyzed reaction of glucose oxidation is as following:

\[
D - \text{glucose} + O_2 \xrightarrow{\text{Glucose Oxidase}} D - \text{gluconolactone} + H_2O_2
\]

A typical decay time-time reaction of optical glucose sensor is shown in Figure 4-11. A typical calibration of \(1/\tau\) versus glucose concentration of this optical glucose sensor is also shown in Figure 4-12.
Figure 4-11: A typical response of glucose sensor to 0.05, 0.1, 0.15 and 0.2 g/L glucose solutions.
Figure 4-12: A typical calibration curve of a glucose optical sensor.
CHAPTER 5 ENERGY METABOLISM OF IVD UNDER COMPRESSIONS

5.1 INTRODUCTORY REMARKS

There are many reasons for people to miss work; one of the leading contributors is low back pain (LBP), which is believed to affect 80% of the population at some point during their lifetime. Suffering from back pain is the chief complaint of 5% of people who visit the doctor in the US (2008). The total medical cost related to low back pain in the US exceeds 100 billion dollars every year (Willis 2009). Many factors were discovered to contribute to LBP. While the exact cause of LBP remains to be clarified, recent studies revealed that intervertebral disc degeneration is closely related to LBP (Luoma, Riihimaki et al. 2000; Roberts 2003). Aging, nutrition supply, mechanical factors and many other aspects contribute to disc degeneration, however, the pathology of disc degeneration is not yet determined (Hadjipavlou, Tzermiadinos et al. 2008; Kandel, Roberts et al. 2008; Adams, Dolan et al. 2009).

IVD is a strong yet deformable tissue that lies between vertebrae and is important for the whole spine structure. Cells only form 1% of the disc volume, however, they play crucial roles in maintaining disc integrity and health by producing ECM components and the chemicals responsible for breaking down the matrix (Bibby, Jones et al. 2001). These processes are highly energy demanding (Im, Freshwater et al. 1976; Hirschberg, Robbins et al. 1998; Prydz and Dalen 2000). Cells consume large amount of adenosine triphosphate (ATP), which is the main cell energy source and generated via glycolysis and mitochondrial respiration with the consumption of glucose and oxygen. IVDs are under daily cycling mechanical loading in vivo; it is reported that mechanical loading can alter transport of oxygen and glucose and release
in the IVD and production of ATP and ECM synthesis of IVD cells *in vitro* (Ohshima, Urban et al. 1995; Huang and Gu 2008; Korecki, Kuo et al. 2009; Fernando, Czamanski et al. 2011; Salvatierra, Yuan et al. 2011). This may indicate the change of overall energy production within the IVD when subjected to mechanical loading. IVD is the biggest avascular cartilaginous tissue in the human body and IVD becomes avascular a few years after the fetal stage, the nutrients supply of IVD is mainly by diffusion, which is not a very efficient transport mechanism and can be hindered by many factors (Urban, Smith et al. 2004); such as endplate calcification and change of proteoglycan content (Grunhagen, Shirazi-Adl et al. 2011). When this transport mechanism is interrupted, the consequence is insufficient nutrient supply which causes decreased energy production and is the main contributing factor in disc degeneration (Urban, Smith et al. 2004; Smith, Nerurkar et al. 2011).

In addition to the important role of intracellular ATP as cellular energy currency, cells constantly release ATP which can regulate cell metabolism, survival, and growth through the purinergic signaling pathway (Burnstock 2006). Hydrolysis of extracellular ATP produces adenosine diphosphate (ADP) and adenosine which can modulate diverse cellular actions via purinergic P2Y and P1 receptors, respectively (Burnstock 2006; Jacobson and Gao 2006), while it also releases inorganic pyrophosphate and phosphate which are strongly associated with mineral crystal formation or tissue calcification (Ryan, Kurup et al. 1992; Johnson and Terkeltaub 2005). Due to the avascular nature of the IVD, ATP and its derivatives may accumulate extracellularly within the disc and be able to influence the biological function of the IVD.

Our recent studies showed that mechanical loading can promote ATP production and release of IVD cells in a 3-dimensional agarose gel model, while cells from two
different anatomical regions, annulus fibrosus (AF) and nucleus pulposus (NP), exhibited different energy metabolism (Fernando, Czamanski et al. 2011; Salvatierra, Yuan et al. 2011). However, the *in vivo* environment involves the cell-ECM interaction, a variety of mechanical events (e.g., tensile/compressive stress and hydrostatic pressure), compression-dependent transport properties, and inhomogeneous distribution of nutrients (Yao and Gu 2007; Huang and Gu 2008). Therefore, the objective of our study is to investigate the energy metabolism in the AF and NP region of the IVD under mechanical loading.

5.2 MATERIALS AND METHODS

Lumbar spines of 4–8 month-old pigs were obtained within 2 hours of sacrifice. Functional spinal units (FSUs) were isolated from lumbar region of the spine by making parallel transverse cuts through the vertebrae. The bone part of the vertebrae was removed till the endplate was disclosed. The FSUs were placed in custom made compression chambers and cultured in high glucose Dulbecco’s Modified Eagle Medium (DMEM, Invitrogen Corp., Carlsbad, CA, USA) containing 10% fetal bovine serum (FBS, Invitrogen Corp, Carlsbad, CA, USA) and 1% antibiotic-antimitotic (Invitrogen Corp, Carlsbad, CA, USA) in an incubator at 37°C overnight. The medium was continuously circulated at 0.37ml/min, which is about twice the rate of blood flow to the disc *in vivo*. For the whole disc compression test, the disc height was measured at 3 locations on the disc surface and then divided in to 5 groups. A custom made bioreactor with circulating culture medium was developed to conduct all compression experiments (Figure 5-1). For static compression groups, the discs were subjected to 10% compressive strain for 2 hours and 1 hour, respectively; for dynamic compression groups, the discs were placed under 1 Hz and 0.1 Hz of sinusoidal compression with 10% maximum strain for 1 hour. The control group was left
undisturbed in the incubator for the duration of the experiment. After the experiment, a transverse cut was made on each unit to expose the AF and NP. The pH value of NP region was detected using pH meter from Cole-Palmar Co. (Vernon Hills, IL, USA). Extracellular ATP concentrations were measured using custom made optical ATP sensor (Wang, Huang et al. 2013). The tissue samples (~10 mm³) were then harvested from the locations shown in Figure 5-2. The samples obtained from AF and NP regions were used to determine the lactate, ATP, and DNA contents. ATP and lactate were released from tissue samples using perchloric acid treatment (Leach 1981) and then determined using the Luciferin-luciferase method (Sigma, St. Louis, MO, USA) and an enzymatic assay (Sigma, St. Louis, MO, USA), respectively.

Cell viability was determined using LIVE/DEAD® Cell Viability Assay (Invitrogen, Carlsbad, CA, USA). DNA content of each harvested sample was determined using a Qubit™ fluorometer and Quant-iT dsDNA HS Assay Kit (Invitrogen Corp., Carlsbad, CA). The lactate and ATP contents were normalized by the DNA content. A one-way ANOVA followed by Tukey test was performed to examine differences in ATP and lactate contents between AF and NP regions and experimental groups of each region using SPSS software (IBM, Armonk, NY, USA).
Figure 5-1: Bioreactor system for whole disc loading experiments
**Figure 5-2:** Harvest locations of AF and NP from IVD

**Figure 5-3:** Actual harvest locations of IVD and porcine IVD dimension
5.3 RESULTS

The live/dead cell staining showed a high cell viability after compression experiments with the AF and NP regions having the cell viability of 88.0%±3.3% and 84.5%±2.5%, respectively (Figure 5-4).

5.3.1 Effects of static compression

Static compression significantly increased lactate accumulation in both AF and NP regions after 1 hour, whereas lactate accumulation was reduced to a non-significant level in the AF and NP regions compared to the non-loading group (Figure 5-5). Accordingly, pH was significantly decreased in both AF and NP regions one hour after application of static compression, whereas after 2 hours of compression, a significant difference in pH was only seen in the AF region, but not in the NP region (Figure 5-6). Total ATP content in all static compression groups was significantly increased (Figure 5-7). In addition, the 1 hour compression group exhibited higher ATP content than the 2 hour compression group for the NP region (Figure 5-7).
Figure 5-4: Cell viability stain of AF (up) and NP (down) tissues. Green indicates live cells while red indicated dead cells.
Figure 5-5: Effects of static loadings on lactate content (n=6), *: significantly different from the control group (p<0.05);
Figure 5-6: Effects of static loadings on pH value (n=8), *: significantly different from the control group (p<0.05).
Figure 5-7: Effects of static loadings on total ATP content (n=8). *: significantly different from the control group (p<0.05); #: significantly different from the other groups (p<0.05).
5.3.2 Effects of dynamic compression

Only dynamic compression of 1 Hz significantly increased lactate accumulation (Figure 5-8). However, all dynamic loading conditions significantly decreased pH and increased total ATP content in both AF and NP regions (Figure 5-9 & 5-10). In addition, frequency-dependent differences were found in lactate accumulation and total ATP content. Lactate accumulation was significantly higher in the 1Hz compression group than the 0.1 Hz and static (0 Hz) compression groups (Figure 5-8) for both AF and NP regions. The total ATP content was significantly higher in the 0.1 Hz compression than the 1 Hz and static (0 Hz) compression groups for the AF region, while the 1 Hz compression group exhibited significantly lower ATP content than the 0.1 Hz and static (0 Hz) compression groups for the NP region (Figure 5-10).

5.3.3 Comparison between AF and NP regions

Under all loading conditions, the NP region exhibited significantly lower pH values and higher total ATP contents than AF (Figure 5-11 and Figure 5-12). There were no significant differences in lactate accumulation between the AF and NP regions for all loading conditions.

5.3.4 Extracellular ATP content in the NP region

A high level of extracellular ATP content was measured at the NP region (165.3±40.8 μM). Under both static and dynamic compression, the extracellular ATP content was significantly reduced in the NP region (Figure 5-13).
**Figure 5-8:** Effects of dynamic loadings on lactate accumulation (n=6). Static compression represents as 0 Hz. *: significantly different from the control group (p<0.05); #: significantly different from the other groups (p<0.05).
Figure 5-9: Effects of dynamic loadings on total ATP (n=8). Static compression represents as 0 Hz. *: significantly different from the control group (p<0.05); #: significantly different from the other groups (p<0.05); & significant difference between two groups (p<0.05).
Figure 5-10: Effects of dynamic loadings on pH value (n=8). *: significantly different from the control group (p<0.05).
Figure 5-11: Comparison of pH between the AF and NP regions (n=7). *: significantly different from the control group (p<0.05).
Figure 5-12: Comparison of total ATP content between the AF and NP regions (n=8). *: significantly different from the control group (p<0.05).
Figure 5-13: Effects of static and dynamic compression on extracellular ATP content in the NP regions (n=5). *: significantly different from the control group (p<0.05); #: significant difference between two groups (p<0.05).
5.4 DISCUSSION

ECM synthesis is an ATP demanding process, especially proteoglycan synthesis in which ATP serves not only as an energy source, but also as a building block in the formation of UDP-sugars and 3’-phosphoadenosine 5’-phosphosulphate (PAPS) (Hirschberg, Robbins et al. 1998; Prydz and Dalen 2000). Because of limited nutrient supply to the IVD, ATP production could be a restrictive factor for maintaining normal ECM synthesis of IVD cells. Furthermore, the IVD is subjected to a variety of mechanical loading, including compression, during spine motion. Compressive loading has been shown to up-regulate ECM gene expression of IVD cells (Chen, Yan et al. 2004; MacLean, Lee et al. 2004; Korecki, Kuo et al. 2009). However, whether up-regulation of gene expressions can increase ECM production may still depend on adequate ATP supply. Although our recent studies showed that both static and dynamic compressions promote energy production of IVD cells in an in-vitro 3-dimensional agarose gel model (Fernando, Czamanski et al. 2011; Salvatierra, Yuan et al. 2011), some in vivo environmental factors were not taken into account, such as cell-ECM interaction and complex mechanical stimuli. Therefore, this study investigated energy metabolism inside the IVD under compression. To our knowledge, this is the first study to demonstrate that mechanical compression affects in-situ energy metabolism in the IVD.

Static compression significantly increased total ATP content and lactate accumulation in both the AF and NP regions (Figure 3), suggesting that static compression promotes glycolysis and consequently increases ATP production in both NP and AF cells. This finding is supported by our previous study which showed that static compression promoted ATP production of AF cells in agarose culture (Fernando, Czamanski et al. 2011). A decrease in solute diffusivity (or tissue
permeability) caused by static compression (Huang and Gu 2008) may also facilitate extracellular lactate accumulation. Furthermore, a decrease in pH caused by static compression in both anatomical regions (Figure 3) may result from the increase in lactate accumulation.

Similarly, 1 Hz dynamic compression significantly increased total ATP content and lactate accumulation in both the AF and NP regions (Figure 4). It is in concurrence with our previous study and suggests that dynamic compression can promote ATP production of AF and NP cells via glycolysis (Fernando, Czamanski et al. 2011). Another factor for increasing ATP production could be an increase in transport of nutrients by dynamic compression (Huang and Gu 2008). In addition, higher lactate accumulation caused by dynamic compression may result in reduction of pH as seen in this study (Figure 4). Furthermore, compared to the other compression groups, significantly higher lactate accumulation found in the 1Hz compression group (Figure 4) suggests that more ATP may be produced in the 1 Hz dynamic compression group. However, significantly lower total ATP content found in 1 Hz dynamic compression group suggests that the 1 Hz dynamic compression group may exhibit higher cellular ATP consumption. These findings also indicate that dynamic compression of 1 Hz promotes energy-demanding cellular activities, such as ECM production (Korecki, Kuo et al. 2009).

Fluid pressurization occurs in the disc, especially in the NP region, when the IVD is subjected to compressive loading (Yao and Gu 2007). Since hydrostatic pressure has been shown to modulate cellular ECM biosynthesis (Ishihara, McNally et al. 1996; Handa, Ishihara et al. 1997), cellular ATP metabolism may be affected by hydrostatic pressure. In the static compression test, when a constant displacement is applied to the IVD, hydrostatic pressure reaches a peak and then gradually reduces to a lower
equilibrium level (Yao and Gu 2007). A statistically significant difference in total ATP content between the one and two-hour static compression groups seen in the NP region could be due to the effect of the time-dependent hydrostatic pressure. In addition, the finding that static compression significantly increased total ATP content in the NP region is not consistent with our previous study (Fernando, Czamanski et al. 2011) which found static compression exhibited no significant effect on ATP production of NP cells in the 3-dimensional agarose model. Since little fluid pressurization occurs in agarose gel under compression due to its high hydraulic permeability (Huang, Hagar et al. 2004), hydrostatic pressure induced in the IVD under static compression is a potential factor causing changes in the total ATP content of the NP region. Furthermore, solute transport and hydrostatic pressure in cartilaginous tissue under dynamic loading depend on loading frequency (Yao and Gu 2004). This could be the reason for the frequency-dependent differences seen in this study. Therefore, the effects of hydrostatic pressure on ATP metabolism of IVD cells needs to be further investigated.

Since both static and dynamic compressions can promote ATP release from IVD cells (Fernando, Czamanski et al. 2011; Salvatierra, Yuan et al. 2011), it is expected that extracellular ATP accumulation occurs in the IVD under compression, especially static compression, which can reduce transport rate of solutes in the IVD (Huang and Gu 2008). However, this study found that both static and dynamic compressions significantly reduced extracellular ATP content in the NP region, suggesting that compressive loading promotes hydrolysis of ATP in the NP region. In addition, high extracellular ATP concentration was detected in the NP region (~ 165 μM). This high extracellular ATP level could be due to the avascular nature of the IVD and the high content of proteoglycans which can inhibit ATP hydrolysis (Vieira, Rocha et al. 2001).
Since extracellular ATP can mediate a wide variety of biological responses via purinergic receptors (Burnstock 2006), a high extracellular ATP level may influence NP metabolism. Furthermore, as described before, compressive loading may promote extracellular ATP hydrolysis, which may result in increases in the extracellular contents of adenine derivatives (i.e., ADP, AMP and adenosine), P Pi, and Pi. Since ADP and adenosine can regulate different cellular activities via P2 and P1 purinergic receptors (Burnstock 2006; Jacobson and Gao 2006), increases in their extracellular concentrations may also have biological effects on NP cells. Due to the capability of P Pi and Pi to regulate crystal formation and tissue calcification (Ryan, Kurup et al. 1992; Johnson and Terkeltaub 2005), increases in the extracellular concentrations of P Pi and Pi in the NP region may contribute to endplate calcification which occurs in aged discs and reduces nutrient supply to the IVD (Grunhagen, Shirazi-Adl et al. 2011). Taken all together, the findings in this study suggest that ATP metabolism may play an important role in maintaining normal biological function of the IVD, while compression-regulated ATP (intracellular and extracellular) metabolism may be a novel mechanobiological pathway for regulating the biological activities of the IVD.

Compared to AF cells, a significantly higher ATP content found in NP cells suggests cellular metabolism in NP cells is more active. This result is consistent with our previous studies (Fernando, Czamanski et al. 2011; Salvatierra, Yuan et al. 2011). In addition, since the concentrations of glucose and oxygen are lower in the NP region than the AF region (Huang and Gu 2008), similar lactate accumulation found in the NP and AF regions suggests that NP cells maintain high ATP content by producing ATP via mitochondrial respiration, which is more efficient in ATP production than glycolysis.

In summary, this study demonstrated that static and dynamic loading increased the
total ATP and lactate contents in the IVD, suggesting that compression can cause an increase in ATP production by promoting glycolysis. Compressive compression can also reduce pH in the IVD due to high lactate accumulation. Due to the capability of ATP to regulate a variety of cellular activities, high extracellular ATP content found in the NP region exhibited and regulated by compression may influence the normal biological function of the IVD. The findings in this study suggest that compression-mediated ATP metabolism could be a novel mechanobiological pathway for regulating the biological activities of the IVD.

However, whether up-regulation of gene expressions can increase ECM production may still depend on adequate ATP supply. Although our recent studies showed that both static and dynamic compressive loading promotes energy production of IVD cells in an in-vitro 3-dimensional agarose gel model (Fernando, Czamanski et al. 2011; Salvatierra, Yuan et al. 2011), the in vivo environmental factors were not taken into account, such as cell-ECM interaction, compression-dependent transport properties, and differences in mechanical stimuli. Therefore, this study investigated energy metabolism inside the IVD under compression. To our knowledge, this is the first study to demonstrate that mechanical compression affects in-situ energy metabolism in the IVD.

Static compressive loading significantly increased total ATP content and lactate accumulation in both AF and NP region (Figure 5-5 & Figure 5-7), suggesting that static compression promoted glycolysis and consequently increased ATP production in both NP and AF cells. This finding is supported by our previous study which showed that static compressive loadings promoted ATP production of AF cells in agarose culture (Fernando, Czamanski et al. 2011). A decrease in solute diffusivity (or tissue permeability) caused by static compression (Huang and Gu 2008) may also
facilitate extracellular lactate accumulation. Furthermore, a decrease in pH found caused by static compression in both anatomical regions (Figure 5-5-Figure 5-7) may result from the increase in lactate accumulation.

Similarly, 1 Hz dynamic loading significantly increased total ATP and lactate accumulation in both AF and NP regions (Figure 5-8 and Figure 5-9). It is in concurrence with our previous study and suggests that dynamic loading can promote ATP production of AF and NP cells via glycolysis (Fernando, Czamanski et al. 2011). Another factor for increasing ATP production could be an increase in transport of nutrients by dynamic loading (Huang and Gu 2008). In addition, higher lactate accumulation caused by dynamic loading may result in reduction in pH as seen in this study (Figure 5-8 and Figure 5-10). Furthermore, compared to the static compression group, significantly higher lactate accumulation found in the 1Hz loading group (Figure 5-8) suggests more ATP was produced in the dynamic loading group. However, no significant difference found in the ATP content between the static and 1 Hz dynamic loading groups suggests that 1 Hz dynamic loading increases ATP consumption. It also indicates that dynamic loading of 1 Hz promotes energy-demanding cellular activities, such as ECM production (Korecki, Kuo et al. 2009).

Fluid pressurization occurs in the disc, especially NP region, when the IVD is subjected to compressive loading (Yao and Gu 2007). Since hydrostatic pressure has been shown to modulate cellular ECM biosynthesis (Ishihara, McNally et al. 1996; Handa, Ishihara et al. 1997), cellular ATP metabolism may be affected by hydrostatic pressure. In the static compression test, when a constant displacement is applied to the IVD, hydrostatic pressure reaches a peak and then gradually reduces to an equilibrium level (Yao and Gu 2007). Significant difference in the ATP content between the one and two-hour static loading groups seen in the NP region could be due to the effect of
the time-dependent hydrostatic pressure. In addition, the finding that static compression significantly increased the ATP content in the NP region is not consistent with our previous study (Fernando, Czamanski et al. 2011) which found static compression exhibited no significant effect on ATP production of NP cells in 3-dimensional agrose model. Since little fluid pressurization occurs in agrose gel under compression due to its high hydraulic permeability (Huang, Hagar et al. 2004), hydrostatic pressure induced in the IVD under static compression may be the main factor causing changes in ATP content of NP region. Furthermore, solute transport and hydrostatic pressure in cartilaginous tissue under dynamic loading depend on loading frequency (Yao and Gu 2004). This could be the reason for the frequency-dependent differences seen in this study. Therefore, the effects of hydrostatic pressure on ATP metabolism of IVD cells needs to be further investigated.

Since both static and dynamic compression can promote ATP release from IVD cells (Fernando, Czamanski et al. 2011; Salvatierra, Yuan et al. 2011), it was expected that extracellular ATP accumulation occurs in the IVD under compression, especially static compression which can reduce transport of solutes in the IVD (Huang and Gu 2008). However, this study found that both static and dynamic loading significantly reduced extracellular ATP content in the NP region, suggesting that compressive loading promotes hydrolysis of ATP in the NP region. In addition, high extracellular ATP concentration is detected in NP region (~165 μM). This high extracellular ATP accumulation could be due to the avascular nature of IVD and the high content of proteoglycans which can inhibit ATP hydrolysis (Vieira, Rocha et al. 2001). Since extracellular ATP can mediate a wide variety of biological responses via purinergic receptors (Burnstock 2006), high extracellular ATP level may influence NP metabolism. Furthermore, as described before, compressive loading may promote
extracellular ATP hydrolysis, which can result in increases in the extracellular contents of adenine derivatives (i.e., ADP, AMP, and adenosine), PPi, and Pi. Since ADP and adenosine can regulate different cellular activities via P2 and P1 purinergic receptors (Burnstock 2006; Jacobson and Gao 2006), increases in their extracellular concentrations may also have biological effects on NP cells. Due to the capability of PPi and Pi to regulate crystal formation and tissue calcification (Ryan, Kurup et al. 1992; Johnson and Terkeltaub 2005), increases in the extracellular concentrations of PPi and Pi in the NP region may contribute to endplate calcification which occurs in aged discs and reduces nutrient supply to the IVD (Grunhagen, Shirazi-Adl et al. 2011). Taken all together, the findings in this study suggest that ATP may play an important role in maintaining normal biological function of the IVD, while compression-regulated ATP (intracellular and extracellular) metabolism may be a novel mechanobiological pathway for regulation of IVD metabolism.

Compared to AF cells, a significantly higher ATP content found in NP cells suggests cellular metabolism in NP cells is more active. It is consistent with our previous studies (Fernando, Czamanski et al. 2011; Salvatierra, Yuan et al. 2011). In addition, since the concentrations of glucose and oxygen are lower in the NP region than the AF region (Huang and Gu 2008), similar lactate accumulation found in the NP and AF regions suggests that NP cells maintain high ATP content by producing ATP via mitochondrial respiration, which is more efficient in ATP production than glycolysis.

In summary, this study demonstrated that static and dynamic loading promoted ATP production and lactate accumulation in the IVD, suggesting that compression can increase in ATP production by promoting glycolysis. Compressive loading can reduce pH in the IVD due to high lactate accumulation. This study also found that the
NP region exhibited high extracellular ATP content which was reduced by compressive loading. The findings suggest that compression-mediated ATP metabolism could be a novel mechanobiological pathway which regulates IVD metabolism.

Low pH was detected in this study especially in NP region after compressive loads (~6.5). It has been shown that low pH can affect many activities of the IVD such as cell viability and protein synthesis rate (Ohshima and Urban 1992; Bibby and Urban 2004). This acidic low pH environment was believed to be fatal for cells (Staub, Baethmann et al. 1990); however, previous studies about IVD cells have shown that decreased cell viability under low pH is also time dependent, which is to say that IVD cells can survive for a short period of time under low pH environment (Liang, Li et al. 2012). Another study also showed that IVD cells cultured in low glucose (0.5 mM) low pH (6.2) conditions for 24 hours have decreased cell viability, from 100% at the beginning to 95% at the end of the experiment (Bibby and Urban 2004). Therefore the low pH (as low as 6.5) condition found in NP region from our study is toxic, but cells can manage to maintain short-term viability under such environment.

A possible limit of this study is that the experimental environment has a different oxygen contents (20%) comparing with in vivo (as low as 1% in central NP region) (Bartels, Fairbank et al. 1998). Cells are believed to utilize glycolysis for the production of ATP under anoxic environments (Romano and Conway 1996). However, glycolysis was observed for IVD cells in vitro which has a culture environment with plenty of oxygen supply (Guehring, Wilde et al. 2009; Fernando, Czamanski et al. 2011; Salvatierra, Yuan et al. 2011). These studies have shown that results obtained from high oxygen contents environments can provide valuable information to explain the responses of cells at in vivo conditions.
5.5 OTHER MEASUREMENTS

Oxygen and glucose distributions in IVDs were also measured using custom-made optical sensors described previously in this dissertation.

A needle type oxygen sensor was inserted from outer AF till reach NP and a caliper was used to record inserting depth. After the measurement the disc was cut transversally and the measurements were corresponded with locations. The result is presented in Figure 5-14. Our finding is consistent with previous researches (Bartels, Fairbank et al. 1998).

Glucose profiles were measured using glucose sensor described in Chapter 4. IVDs were cultured in high glucose (4.5 g/L D-glucose) and low glucose (1 g/L D-glucose) Dulbecco’s Modified Eagle Medium (DMEM, Invitrogen Corp., Carlsbad, CA, USA) containing 10% fetal bovine serum (FBS, Invitrogen Corp, Carlsbad, CA, USA) and 1% antibiotic-antimitotic (Invitrogen Corp, Carlsbad, CA, USA). No significant difference was found in glucose content between the measurement locations in the AF and NP regions when the IVD was cultured in high glucose DMEM. However, AF had higher glucose concentration than NP when low glucose DMEM was used for disc culture (Figure 5-15). This measurement is consistent with previous study (Jackson, Huang et al. 2011).
Figure 5-14: Oxygen profiles measured in the IVD using a needle type optical oxygen sensor.
Figure 5-15: Glucose contents of AF and NP under low glucose DMEM culture.
CHAPTER 6 CONCLUSION AND RECOMMENDATIONS

6.1 SUMMARY AND CONCLUDING REMARKS

Low back pain is a big social burden which affects 80% of people in USA at certain time point during their lives and cost numerous societal resources. Each year more than 50 billion US dollars are spent for low back pain in United States (Borenstein, Calin et al. 2012). Although it is not very well studied yet, intervertebral disc degeneration was revealed to be a possible cause of low back pain (Cox 2011). IVD is the biggest avascular tissue in human body, the pathway of nutrient supply is through peripheral blood vessels. For NP cells the distance to closest blood vessel can be as far as 20mm, which makes the environment of NP cells one of the worst in human body with regards to having a very poor nutrient supply (Grunhagen, Shirazi-Adl et al. 2011). Cells play pivotal roles in IVDs by synthesizing and breaking down ECM to maintain a favorable environment for tissue functions, however these activities are high energy commanding (Urban and Roberts 2003; Bibby, Jones et al. 2005). Thus nutrient supply can be a limit factor for cells to maintain ECM integrity. Previous researches showed that mechanical loadings can affect cell energy metabolism (Lee, Wilkins et al. 2002; Fernando, Czamanski et al. 2011; Salvatierra, Yuan et al. 2011). However, there are more mechanical events inside IVDs when mechanical loadings are applied, such as hydrostatic pressure, cell-ECM interaction, etc. All of these have been showed to affect ECM synthesis of IVD cells (Ishihara, McNally et al. 1996; Setton and Chen 2006; Korecki, Kuo et al. 2009). Therefore it is essential to study the energy metabolism of IVD under mechanical loadings, especially compression which is the most common mechanical loading.
Hence, the main objectives of this dissertation were threefold: (1) To design and build a novel bioreactor system for whole IVD disc culture and perform static and dynamic compression studies; (2) To develop novel optical sensors to detect glucose and ATP contents and (3) To investigate the effects of static and dynamic compressions on energy metabolism of IVD. Three studies were carried out to achieve the following specific aims: (1) A bioreactor system were built to perform both static and dynamic compressive loadings with attached culture medium circulation and loading force detecting and recording devices (Chapter 3); (2) An optical fiber based ATP sensor was fabricated using sol-gel entrapping enzyme technique (Chapter 4) and (3) the changes on lactate accumulation, pH value, total ATP and extracellular ATP were measured in the porcine IVD under static and dynamic compression. Measurements of oxygen tension and glucose distribution were also performed (Chapter 5). The important findings from these studies are discussed as following.

6.1.1 Optical fiber based ATP sensor

A novel optical fiber based ATP sensor was fabricated. Two layers of sol-gel coating layers were applied to the end of optical fiber, the first layer contained Ruthenium complex which can be excited by blue light (465 nm wave length) and emits red light (610 nm wave length); this red light can be absorbed by oxygen molecules without radiation, thus quench the signal and increase decay time ($\tau$).
The second layer entraps two enzymes: glycerol kinase and glycerol 3-phosphate, which can catalyze the oxidization of ATP and consume oxygen. Reverse of decay time (1/τ) has a linear relationship with oxygen concentration. Therefore ATP content can be detected by measuring the change of decay time.

This sensor has a broad detecting range from $10^{-3}$ mM to 1.5 mM and is not affected by physiological pH value change or ATP derivatives such as ADP, AMP and adenosine (Wang, Huang et al. 2013). Since this sensor measures the change of oxygen partial pressure at the local micro-environment of coating layers, we also developed a compensation method for different environmental oxygen contents.

Oxygen sensor was fabricated by coating the first layer only as subscribed above; glucose sensor was developed by the same method, only replace the enzymes glycerol kinase and glycerol 3-phosphate to glucose oxidase. Oxygen sensors and glucose sensors all have robust performance and are suitable for physiological tissue in situ measurements.

6.1.2 Effects of static and dynamic compressions on IVD energy metabolism

Five (5) groups of static and dynamic compressive loadings were performed: control group without any mechanical loadings (0 Hz 0 hour group), 1 hour static compression group (0 Hz 1 hour group), 2 hour static compression group (0 Hz 2 hour group), 1 hour 0.1 Hz group and 1 hour 1 Hz group. 10% strain was applied to compressive groups.

A transversal cut was applied to the disc after experiment; lactate accumulation and total ATP contents were measured using corresponding assays. Extracellular ATP
contents at NP region were detected using ATP sensors described earlier (Wang, Huang et al. 2013). Oxygen distribution inside of IVD and glucose contents at AF and NP regions were also measured using custom made sensors.

We discovered that both static and dynamic loadings tend to increase lactate accumulation in AF and NP regions; and significantly decrease the pH value; total ATP contents were significantly increased in both AF and NP regions, these suggests that mechanical loadings promote ATP production via glycolysis which produces lactate, causes the decrease in pH value. However, we also detected significant decrease of extracellular ATP in NP region, this may due to the increase of ATP hydrolysis induced by mechanical loadings. The measured high extracellular ATP contents in NP region (~160 μM) also reveal certain pivotal environmental condition for NP cells.

In summary, this study helps us to better understand the energy metabolism of IVD under compressive mechanical loadings, which is essential to better understand disc degeneration and low back pain and may also be beneficial for future endeavor on disc regeneration.

6.2 RECOMMENDATION FOR FUTURE STUDIES

The overall objective of this study was to elucidate energy metabolism of IVD under compressive loading. Although valuable insights of energy metabolism changes induced by compressions were provided in this dissertation, there are still many important works left to finish under this big frame. Therefore recommendations of future work are described as following. Compression experiments reveals that lactate accumulation in both AF and NP region were promoted by compressive loadings and cause a significant
decrease in pH (to as low as 6.5). This creates a harsh environment for IVD cells, especially NP cells, which are surrounded by dense AF tissue and up to 20 mm away from nearest blood supply (Moore 2006). How IVD cells survive and maintain activities under such difficult conditions remains an enigma. Therefore a study about IVD cell response and activities at low pH environment should be conducted.

It was also found that NP region maintains high extracellular ATP contents (~160 μM); this may reveal a vital condition for NP cells to survive in the harsh mechanical and biochemical conditions and maintain their normal biological function (e.g., ECM biosynthesis) and notochordal phenotype. Effects of high extracellular ATP on NP cells could also be studied. Our study showed that extracellular ATP contents were significantly decreased for all compression groups. This indicates that compressions may promote ATP hydrolysis and thus increase local concentrations of adenine derivatives (i.e., ADP, AMP and adenosine) and phosphate. This should be elucidated by future works on measuring changes in the contents of ADP, AMP and adenosine of NP region under compressive loading. Calcium has a high concentration in cartilage but is mostly bound in proteoglycan as part of the anionic groups and hence not available for precipitation. Phosphate can replace calcium by a ion-exchange effect and thus increase endplate calcification (Hunter 1987). Therefore the relation of compressive loadings on IVD, hydrolysis of ATP and endplate calcification should be further studied. Other products of ATP hydrolysis include ADP, AMP and adenosine.

Adenosine may also play a very important role on protecting notochordal cells which was reported to protect the NP from degradation (Erwin, Islam et al. 2011). A future
study should be carried out to investigate the effects of adenosine on preventing IVD cells degeneration.

A discrepancy between this study and previous agarose model study about effects of static compression on IVD ATP production was mentioned in Chapter 5. As discussed in that chapter, this may due to the hydrostatic pressure inside IVD tissue. Thus a future study should be conducted to investigate the effect of hydrostatic pressure on ATP metabolism of IVD cells, especially on NP cells which undergo high hydrostatic pressure during static and dynamic compressions.

Glycolysis produces lactate and less ATP than mitochondrial respiration and is likely the dominant energy production method for NP which has limited oxygen supply. It is essential to solve the problem of nutrition supply for NP cells to acquire enough energy to maintain extracellular matrix integrity, which is essential to prevent degradation of IVDs (Urban and Roberts 2003). Inability of NP to maintain integrity of ECM will cause the IVDs to lose their mechanical properties and further lead to disc degeneration (Roughley 2004). Therefore, works should be carried out in investigating the effect of promoting nutrient supply to NP on retarding or even inverting disc degeneration.

In summary, the recommended studies would extend the accomplishment of the studies in this dissertation and help to achieve the long term goal which is to elucidate the pathophysiology of disc degeneration and low back pain, and develop treatment to retard or repair disc degeneration and cure low back pain.
REFERENCES


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