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The Impact of Oxidative Stress on Postmortem Meat Quality

by

Nicolas J. Herrera

A Thesis

Presented to the Faculty of
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THE IMPACT OF OXIDATIVE STRESS ON POSTMORTEM MEAT QUALITY

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University of Nebraska, 2020

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Tenderness is the primary factor to influence consumer palatability in fresh meat. The tenderization of muscle is dependent on proteolytic mechanisms, including calcium-dependent, non-lysosomal endogenous enzymes such as calpains and proteasomes. These mechanisms have been indicated alongside apoptosis early postmortem as cascades of apoptotic events promote ideal conditions for postmortem tenderization. Recent literature has identified oxidative stress-mediated events to be linked to apoptotic activity. Oxidative stress is defined as the overwhelming production of reactive species compared to homeostatic, endogenous antioxidants present within muscle systems. These conditions exhaust antioxidant function and facilitate reactive species to alter protein, lipid, or nucleic morphology and functionality. Perhaps, the generation of oxidative stress can alter muscle tissue, resulting in changes in the overall quality of fresh meat postmortem. This study examined the impact of different levels of oxidative stress *in vivo* on postmortem muscle quality of lambs, with emphasis towards elements of proteolytic mechanisms responsible for meat tenderness. Lipopolysaccharides (LPS) are endotoxins used to induce acute stress for this experiment. Effects of inducing lambs with an injection of either a saline control, 50 ng of LPS per kg of bodyweight (LPS50), or 100 ng of LPS per kg of bodyweight (LPS100). Injections were administered every 72 hours

across a nine-day immune challenge. Treatments used examined changes in biomarkers of oxidative stress (RNA Transcriptomics and isoprostanes), pre-harvest stress conditions (rectal temperature), free calcium concentration, proximate composition, color and lipid oxidation, and tenderness. In this study, lambs administered LPS50 tended to be more tender early postmortem, and had a greater degree of troponin T degradation compared to control samples ($P < 0.05$). The LPS-treated lambs exhibited noticeable upregulation of gene pathways responsible for cell growth, differentiation, degradation, and generation of oxidative species and antioxidants ($P_{\text{raw}} < 0.05$). While not significant, LPS treated samples exhibited more oxidative biomarkers present in muscle tissue (isoprostane content). The LPS treatments had no detrimental effects of color or lipid oxidation ($P > 0.05$). Oxidative stress may impact early postmortem tenderization of meat.

Key words: apoptosis, lamb, meat quality, oxidative stress, proteolysis, tenderization

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INTRODUCTION

This study seeks to understand the effect of oxidative stress *in vivo* on post-harvest changes in meat quality which may improve tenderness early postmortem. Tenderness is repeatedly cited as the primary element associated with both eating quality and consumer purchasing decisions (Miller et al., 2001; Platter et al., 2005). In the U.S., consumers are willing to pay premiums for higher quality meat to ensure a good eating experience (Platter et al., 2005). However, noticeable variation exists when comparing tenderness (shear force values), in particular when comparing across marbling scores in beef. More so, the extent of variation in tenderness increases as marbling decreases. Inconsistent meat tenderness and its impact on consumer satisfaction is an obstacle to optimizing both domestic and international demand for U.S. meat products. Thus, investigations into the process of postmortem tenderization and the role of cellular organelles and mechanisms involved have strong, practical application.

Muscle contraction is the result of a complex mechanism, requiring multiple proteins and molecular compounds to work in concert. Contraction of myofibrillar proteins allows for the physical movement of bones and tendons in response to neural stimuli (Kuo et al., 2015). The postmortem deterioration of these proteins alters meat tenderness. Further, if these protein complexes are altered *in vivo*, an earlier onset of the tenderization process might be achieved, promoting more tender beef. Using proteomics approaches, Malheiros et al. (2019) analyzed beef muscle tissue across samples distinguished as tough, intermediate, or tender based on their average shear force values (kilograms-force) and identified different oxidized proteins across each tenderness group. The tender group samples, compared to the tough and intermediate groups, had highly

oxidized structural, contractile, and regulatory proteins, all directly associated with muscle contraction and tenderization mechanisms. Concurrently, the tender group exhibited higher oxidized proteins related to enzyme regulation (peroxiredoxin; superoxide dismutase [Cu-Zn]) and cell protection (heat shock proteins). From these results, it is clear that tender muscles exhibited oxidation of specific proteins associated with meat tenderization. There is tremendous value to further understanding why certain proteins oxidize in a manner to promote more tender beef.

Predisposition to oxidative stress may promote an increase in oxidized proteins. In mitochondria, the oxidative phosphorylation mechanism produces reactive oxygen species (ROS), the driving force of oxidative damage. These are highly reactive free radical compounds generated as by-products of ATP synthesis, such as the production of superoxide anion (O_2^-) and hydrogen peroxide (H_2O_2) (Sierra et al., 2013). Due to their unstable state, ROS will collect electrons from their immediate surroundings, commonly from organelles, to reach a stable state. The interaction of ROS with components of the muscle can result in major cell damage, lipid oxidation, protein turnover, and DNA alteration (Scicchitano et al., 2018). To alleviate this damage, muscle cells contain various non-enzymatic and enzymatic antioxidants to detoxify ROS and prevent cell damage (Kozakowska et al., 2015). These antioxidant mediators include superoxide dismutase (SOD), catalase (CAT), glutathione peroxidase (GPX), and other low molecular free radical scavengers (Bekhit et al., 2013; Scicchitano et al., 2018). The dietary antioxidant alpha-tocopherol, vitamin E, has also been shown as an effective supplement to alleviate ROS generation (Harris et al., 2001; Maraba et al., 2018).

However, under conditions of significant oxidative stress, endogenous antioxidants may not be sufficient to quench the increased production of free radicals.

In states of prolonged oxidative stress, stability of mitochondrial membranes will fail, thereby allowing encapsulated ROS generated by the mitochondria to be released to interact with the matrix of muscle cells (Paradies et al., 2001; Elmore et al., 2007; Powers et al., 2011). From this phenomenon, there is growing evidence that oxidative stress can promote programmed cell death (apoptosis) and enhance early postmortem proteolysis. Oxidative stress during death allows the release of cytochrome c, a pro-apoptotic heme protein localized between the inner and outer mitochondrial membranes (Cai et al., 1998). The cascade of the apoptotic pathway gives rise to the activation of caspases, allowing subsequent proteolytic enzymes such as calpains (μ and m) and cathepsins to interact with the muscle tissue, promoting structural degradation of muscle proteins during postmortem aging (Logue et al., 2008; Kemp et al., 2010). Wang et al. (2018) investigated the influence of ROS-generated oxidative stress in pre-rigor muscle tissue. They concluded that increased generation of oxidative stress via ROS accumulation prior to animal harvest may promote apoptotic factors such as cytochrome c displacement and the caspase cascade. They also speculated that these conditions could initiate proteolytic degradation of muscle fibers and improve meat tenderness. The influence of oxidative stress is also being investigated for its impact on other factors of meat quality, such as lipid oxidation and color stability.

The working hypothesis is that controlled levels of induced oxidative stress in lamb can stimulate the production of ROS, generating a state of oxidative stress that modifies the function of the different mechanisms responsible for meat quality. If the

hypothesis is true, control of oxidative stress-inducing events for the live animal may impact meat quality.

The objectives of the research were to understand the mechanisms and components related to meat quality in lamb from wethers administered defined levels of an oxidative stress promoter. Specific objectives were to:

- 1) Identify changes in physiological status of lambs via a lipopolysaccharide (LPS) challenge.
- 2) Characterize the relationship between muscle quality and oxidative stress on the basis of the RNA transcriptome as it relates to LPS and Control-treated lambs.
- 3) Quantify changes in oxidative biomarker generation in relation to LPS-challenged lambs.
- 4) Assess the effects of oxidative stress parameters on lamb longissimus muscle tenderness; and
- 5) Evaluate the impact of known levels of oxidative stress on color stability of intact fresh lamb meat.

The long-term goal of this research is to improve the tenderness of fresh meat by understanding the complexity of muscle tenderization as affected by *in vivo* oxidative stress. The results are intended to serve as a model for future analyses in beef, meant to understand the impact which oxidative stress-induced conditions may change the overall meat quality attributes. This research could provide insightful thought into how biochemical mechanisms are altered due to inherent physiological stress that occur within

our current livestock strategies, and how these strategies could be manipulated to optimize the greatest opportunity for more consistently tender meat.

LITERATURE REVIEW

Mechanism of muscle contraction

Muscle contraction is the result of a complex mechanism, which requires multiple proteins and molecular compounds to work in concert (Kuo and Ehrlich, 2015). In the event of a stimuli, a nerve response instigates signal movement to the neuromuscular junction surrounding the muscle fiber. This results in the binding of acetylcholine to facilitate transfer of sodium ions through the sarcolemma, allowing an action potential to travel down the t-tubule. As a result, an opening of calcium ion channels allows the sarcoplasmic reticulum (SR) to release calcium ions via the ryanodine receptors. The main function of calcium during muscle contraction is to bind to troponin C, one of the three troponin subunits of the regulatory protein that facilitates tropomyosin to expose or cover actin binding sites. When exposed, myosin hydrolyzes adenosine triphosphate (ATP) to trigger the power stroke, pulling the actin filament towards the M-line, shortening sarcomere length and completing contraction. Subsequent ATP binding is required to release myosin heads and decrease the degree of contraction. This implies muscle contraction is dependent on the relationship between the efficient calcium regulation of the SR and ATP generation within the mitochondria (Dirksen, 2009).

Sarcoplasmic reticulum regulatory proteins

The intracellular signal for muscle contraction, regulation of calcium storage, release, and cytoplasmic uptake are all factors manipulated by the SR. Storage of calcium ions is met by homeostatic control of luminal calcium binding proteins. Calsequestrin, the most abundant luminal calcium binding protein in the SR, maintains a high binding

capacity to support a large concentration of calcium ions reserves within the SR, while also allowing a rapid release of calcium ions up reaction to a stimulus via an action potential (Rossi and Dirksen, 2006). Calsequestrin also is bound to the SR membrane via interactions with ryanodine receptors (RyR1), facilitating effective calcium release. Activity of RyR1 is mutually dependent on bound calsequestrin, as an interrelationship between levels of total luminal calcium can influence binding capacity of calsequestrin (Zhang et al., 1997).

Calcium release is controlled by the direct protein-protein interactions as a result of sodium-ion transfer with RyR1 proteins and inositol triphosphate (Meissner, 2002). A cysteine-dense protein matrix, RyR1 is the predominant protein responsible for the release of calcium from within the SR of skeletal muscle (MacKrell, 2012). Regulation of RyR1 is dependent on calcium concentration (μM to mM) within and surrounding the SR, as high levels within the sarcoplasm inhibit calcium release and low levels promote calcium release (Copello et al., 1997). As an action potential travels down the t-tubule, excitation coupling reactions take place, promoting a conformational change to the dihydropyridine receptor (DHPR) and initiating the activation of RyR1 (Rossi et al., 2009). In conjunction with calsequestrin, activity of RyR1 is manipulated by free radical-mediated changes in amino acid structure during states of physiological stress, illustrating the relationship between proteins (Eu et al., 1999). Inositol triphosphate acts as signaling molecule responsible for calcium release along the SR. Inositol triphosphate assists in prolong levels of calcium in the sarcoplasm upon RyR1 release.

Calcium influx is achieved by the Sarco-endoplasmic reticulum ATP-ase pumps (SERCA). Sarco-endoplasmic reticulum ATP-ase is comprised of two primary structures:

a transmembrane portion that facilitates opening and closing of SERCA pumps and a protein head which sustains calcium binding and release (Toyoshima and Inesi, 2004). As its name suggests, SERCA is an energy-dependent protein pump which requires the hydrolysis of adenosine triphosphate (ATP) (Rossi and Dirksen, 2006). Within muscle, two isoforms are present, with SERCA 1 and SERCA 2 present in the sarcolemma of Type II and slow-twitch skeletal muscle fibers, respectively (Ishii et al., 1998). Similar to RyR1, SERCA activity can be inactivated in the presence of high free-radical concentration and inhibited by sarcolipin (Ishii et al., 1998). All these proteins are located along the outer membrane of the SR and each protein is involved with the complex action of calcium flux during muscle contraction and relaxation.

Membrane composition

Organelle membranes, focusing on SR and mitochondria, are generally comprised of a phospholipid bilayer, with hydrophobic fatty acid tails oriented towards the interior of the membrane and the hydrophilic fatty acid head facing towards the cytosol exterior, allowing an impermeable matrix to prevent transfer of water-soluble molecules (Borchman et al., 1999). Composition of fatty acids is critical to fluidity and function of bound protein channels (Stanley and Parkin, 1991). Saturated fatty acids contain no double bonds across the carbon chain, lowering overall fluidity due to their densely packed spacing. Conversely, unsaturated fatty acids contain one or more double bonds across their hydrocarbon chain, allowing intermittent spacing across the membrane to increasing fluidity (Borchman et al., 1999). Under states of oxidation, the fatty acid composition is paramount to influence downstream effects on meat quality (Faustman et

al., 2010). Allylic carbons found on unsaturated fatty acids have a weaker capacity to retain their hydrogens compared to saturated fatty acids, allowing an affinity for greater oxidation at these sites, altering their physical orientation. This initiates a cascade of oxidizing reactions, impacting the functionality of *in vivo* organelle membranes and postmortem attributes of meat quality such as color stability, lipid oxidation, and meat tenderization.

Generation of oxidative properties

Oxygen comprises roughly 21% of atmospheric air, nearly all of which is utilized by mitochondria during respiration (Bolisetty and Jaimes, 2013). After binding to hemoglobin, oxygen is transferred to myoglobin with muscle cells and available to perform oxidative phosphorylation within mitochondria. This mechanism acts as the production cycle of ATP, the essential energy source of biological function. Constant metabolism of oxygen is necessary in order to supply sufficient ATP for basal metabolic and physical function within an organism. While a necessity to sustain life of aerobic organisms, such as livestock, oxidative phosphorylation does not come without byproducts. These byproducts are beneficial in some respect for cellular signaling and inflammatory response, but an intricate balance of these molecules is needed to prevent detrimental quantities that can alter cellular stability. It is critical, therefore, to further understand the generation of these products in conjunction with mitochondrial function and their potential impact on muscle cells.

Mitochondrial function in skeletal muscle

Mitochondria are integral to normal cellular function and are the key contributor to energy production in eukaryotic cells. Originating as a specialized form of bacteria, mitochondria have adapted an endosymbiotic relationship within eukaryotic cells, acting as the primary facilitator of phospholipid and heme synthesis, calcium homeostasis, apoptotic activation and cellular death (Gray et al., 2001; Bolisetty and Jaimes, 2013). Mitochondria hold key enzymatic systems designated for the oxidation of carbohydrates, fat, and proteins to produce cellular energy in the form of ATP. The production of ATP is essential for maintenance of skeletal muscle function, including contraction, relaxation, signal transduction, and energy-dependent enzymes utilized in molecular reactions. Mitochondria are comprised of four distinct domains, each holding their unique function in relation to skeletal muscle (Kühlbrandt, 2015).

Mitochondria structure

Commonly misinterpreted as oval shaped organelles, mitochondria are very diverse in their shape due to cycles of fissions and fusions with other mitochondria to better facilitate the demands of cellular respiration (Gray et al., 2001). Given the current state of an organism or localized cells, mitochondrial fission or fusion is regulated to optimize energy metabolism while maintaining a homeostatic function within the mitochondrial membrane. This is shown when overproduction of H_2O_2 facilitates fission to repair damaged mitochondrial matrices (Schenkel and Bakovic, 2014). Focusing on mitochondrial structure, the outer membrane is comprised of a phospholipid membrane and separates the organelle from the cytoplasm. Porous in its design, the outer membrane

facilitates transportation of small uncharged molecules (<5 kDa) via the voltage dependent anion channel (VDAC). The inter-membrane space allows translocation of ions further into the mitochondria. In contrast to the outer membrane, the inner membrane is a tightly built diffusion barrier, designed for selective ion and molecular transport. This exclusivity is shown by an inner membrane potential of approximately 180mV, illustrating the movement of ions via specialized transport proteins (Schenkel and Bakovic, 2014). Next, the inner mitochondrial membrane is distinguished as two subsections, the boundary membrane and the cristae. As closely stacked discus-shaped structures, cristae are bound to the majority of the inner membrane, and act as the site of mitochondrial energy conversion (Osellame et al., 2012; Kühlbrandt, 2015). Lastly, the inner mitochondrial matrix is the cytoplasmic space responsible for precursor mechanisms in oxidative phosphorylation and mitochondrial DNA (mtDNA) synthesis. This is possible due to a higher pH found within the mitochondrial matrix, permitting regulation of ions via an electrochemical gradient during oxidative phosphorylation (Llopis et al., 1998).

Mitochondria uniporter

During homeostasis, high concentrations of extracellular calcium are capable of entering the mitochondria via the mitochondria uniporter. Due to the close proximity of mitochondria to SR organelles, evolutionary development of this protein was necessary (Dirksen, 2009). This protein regulates the passive uptake of calcium while maintaining an even distribution of calcium concentration across the surrounding cytosol and the mitochondria. Acting as a buffer for calcium in the cell, the uniporter allows the

concentration of calcium in the mitochondria to reach 10^6 greater concentration compared to the cytosol. The concentration gradient of calcium acts to maintain proper membrane potential of mitochondria, which is essential for proper function the TCA cycle and function of the electron transport chain (Santo-Domingo and Demaurex, 2010). Given the low affinity of the uniporter, however, a large amount of calcium is required to activate the uniporter to cycle calcium into the mitochondrial matrix (Kirichok et al., 2004).

Mitochondria permeable transition pore

During mitochondrial storage of calcium, there are methods to mitigate calcium overload. The mitochondria permeable transition pore (MPTP) is a secondary method to release mass quantities of calcium in case of possible mitophagy. Used as a final option, the MPTP opens the matrix to release large quantities of calcium as quickly as possible during state of mitochondrial stress. However, calcium discharging can prolong exposure of pore openings, facilitating disruption of the mitochondrial matrix's membrane potential, disrupting the mitochondria's capacity to transfer electron and protons across the matrix and lead to cellular death (Di Lisa et al., 2001).

ATP production precursors

Oxidative phosphorylation is the mechanism by which nutrients are oxidized to produce electrons, enabling the generation of ATP (Santo-Domingo and Demaurex, 2010; Osellame et al., 2012). The subsequent breakdown of proteins, fats, and carbohydrates produce the end product acetyl-CoA via amino acid metabolism, β -oxidation, and glycolysis, respectively. Dependent on the nutrient, each pathway provides

a unique technique for acetyl-CoA to enter the mitochondrial matrix (Owen et al., 1998). Amino acid metabolism degrades amino acids to a variety of intermediates for the citric acid cycle and/or pyruvate for gluconeogenesis. During a fasting state, oxidation of amino acids predominantly occurs when muscle cells break down amino acid constituents with assistance of the liver, with some additional metabolism in the kidneys and small intestine (Owen et al., 1998). The end product, pyruvate, is subsequently brought into the mitochondrial matrix from the cytosol via the mitochondrial pyruvate carrier. Beta-oxidation of lipids is a more complex pathway, requiring the activation of acyl-CoA synthetase to form fatty acyl-CoA. This facilitates fatty acids to pass through the outer mitochondria membrane via the carnitine palmitoyl transferase I (CPTI). When reacted with carnitine, a component of amino acids, acyl-CoA forms acyl-carnitine, and is then allowed to pass into the inner mitochondrial membrane via translocase, with byproduct CoA-SH is sent back into the cytosol (Melzer, 2011; Dunning et al., 2014). Carbohydrate metabolism is fairly similar to amino acid metabolism in its end product. After the breakdown of simple sugars glucose, fructose, and galactose via glycolysis, pyruvate can be transported into the mitochondrial matrix via the mitochondrial pyruvate carrier (Owen et al., 1998; Melzer, 2011). Carbohydrate breakdown is a result of each sugars chemical makeup and how metabolic organs facilitate degradation. To utilize fructose, the liver is the predominant organ needed to convert to phosphorylation substrates, as fructose is not as readily absorbed in the blood as glucose. This constraint requires more complex degradation techniques (Melzer, 2011). Once nutrients are broken down into substrates, oxidative phosphorylation can occur.

Oxidative phosphorylation

During oxidative phosphorylation, a choreographed exchange of electron donors works in concert with respired oxygen and hydrogen ions to produce ATP molecules. Mitochondria bioenergetics facilitate electron donors nicotinamide adenine dinucleotide (NADH) and flavin adenine dinucleotide (FADH₂) via the citric acid cycle (Osellame et al., 2012). Matrix protein complexes (MPC I, III, IV) embedded in the inner membrane to facilitate the pumping of protons from the inner mitochondrial matrix into the inner membrane space. These complexes work in conjunction with MPC II, coenzyme Q, and cytochrome c to continue the transfer of electrons within the inner membrane. This alters the membrane potential, increasing an affinity for hydrogen ions to re-enter the mitochondrial matrix. Appropriately, MPC V permits transfer of hydrogen ions back into the mitochondrial matrix, and when bound with enzyme ATP synthase, it is this transfer of ion that facilitate ATP generation from previous oxidized adenosine diphosphate (ADP) (Osellame et al., 2012; Kühlbrandt, 2015). This mechanism acts as the primary pathway towards energy production and produces multiple byproducts as a result. While major energy donors NAD⁺ and FADH⁺ are recycled in the mitochondrial matrix to regenerate to their reduced state, the production of additional, more unstable molecules, can serve further biological functions.

Reactive oxygen species

Free radicals are necessary byproducts of oxidative phosphorylation. They are developed and controlled within the mitochondria. While free radical formation is commonly associated with states of inflammation or chronic illness, the vast majority of

free radical species are formed during basal physiological functions of cells, such as oxidative phosphorylation (Dröge, 2002; Bolisetty and Jaimes, 2013). The exchange of electrons from NADH and FADH₂, coupled with the transfer of H⁺ ions through the inner mitochondrial membrane, facilitate the generation of oxygen-based radicals known as reactive oxygen species (ROS).

Other components of muscle cells (sarcoplasmic reticulum; sarcolemma) have been suspected to contributing to ROS production, primarily due to subtle interactions with cofactor NADPH and membrane proteins (Xia et al., 2003). However, the mitochondria are widely regarded as the central point of ROS generation. Reactive oxygen species include negatively charged, oxygen centered radicals (O₂⁻, OH⁻), but also include reactive derivatives such as hydrogen peroxide (H₂O₂) (Zorov et al., 2000; Dröge, 2002; Powers et al., 2011; Xing et al., 2019). Low physiological levels of oxidants and other radicals play an important role in the control of gene expression and regulation of cell signaling. As signaling transduction molecules, ROS can influence redox-sensitive pathways in skeletal muscle to alter how proteins modulate growth, differentiation, proliferation, and controlled muscle turnover. Concurrently, ROS levels can influence redox-sensitive pathways such as gene expression in response to inflammation, exercise, infection, growth factors, and stress. In particular, the mitogen activated protein-kinase (MAPK), nuclear factor kappa-light chain enhancer of activated B cells (Nf-kB), and activator protein 1 (AP1) have all be shown to change their expression in response to ROS (Powers et al., 2011). The generation of ROS, when coupled with minimal physiological response, holds potential to alter the composition of cellular stability through oxidative stress and change myocyte composition.

Superoxide

Superoxide (O_2^-) is the primary ROS generated by incomplete reduction of oxygen in the electron transport chain *in vitro*, and is an intermittent product of specific enzymatic systems. Through comprehensive review, it is theorized that the majority of O_2^- production occurs within the mitochondrial matrix during the electron exchange at Complex I (Ott et al., 2007; Bekhit et al., 2013). This phenomenon is stimulated with the presence of succinate, a common substrate produced from Complex III. While Complex III can generate O_2^- , inhibition of Complex III via antimycin reduces its activity along the electron transport chain, which is theorized to significantly increase superoxide generation. Because antimycin does not inhibit Complex I, it can continue to utilize electron donors and generate O_2^- (Ott et al., 2007; Powers et al., 2011; Bolisetty and Jaimes, 2013). In contrast to Complex I, O_2^- production at Complex III is possible outside of the mitochondrial matrix within the inter-membrane space. Coenzyme Q10, (i.e. ubiquinone) is a lipophilic electron carrier that is conjoined to Complex III (Paradies et al., 2001). Through a series of redox reactions, Coenzyme Q10, ubiquinone, is capable of transferring electrons across Complex III. As a result, the recycling oxidation-reduction reactions of ubiquinone also facilitate the generation of O_2^- (Ott et al., 2007; Bolisetty and Jaimes, 2013). While considered relatively unreactive in comparison to other radicals, superoxide can extract electrons from biological membranes and other cellular components. An example of this is the reduction of cytochrome c to initiate apoptotic mechanisms. Superoxide does, however, hold a noticeably longer half-life (~5 seconds) than most radicals, and is capable of dismutating with other molecules, such as nitric oxide (NO^-) and hydrogen ions (H^+) to produce ROS which are more unstable than O_2^-

(Zorov et al., 2000; Fulle et al., 2004). While superoxide is constantly produced at low levels during normal respiration, intracellular reactions permit superoxide to have an increased impact on organelle and cellular function.

Hydrogen peroxide

Hydrogen peroxide (H_2O_2) is produced by O_2^- binding with hydrogen ions or enzymatic breakdown of O_2^- via manganese-superoxide dismutase enzymes. Organelle membranes are permeable to H_2O_2 , allowing it to cross the mitochondria into the cytosol with the use of aquaporins surrounding the outer mitochondrial membrane (Bienert et al., 2006; Bolisetty and Jaimes, 2013). While it is speculated O_2^- may also diffuse through mitochondria by voltage dependent anion channels, it is unknown to what degree this diffusion occurs (Madesh and Hajnóczky, 2001). As a non-radical ROS, H_2O_2 is incapable of directly oxidizing lipids or DNA, but it can be cytotoxic during periods of chronic illness such as cancer (Fulle et al., 2004; Powers et al., 2011). Hydrogen peroxide can act as a major precursor to oxidative damage within a cell. In particular, when H_2O_2 is converted into hydroxyl radicals (OH^-) in the presence of ferrous (Fe^{2+}) and cuprous (Fe^+) iron via the Fenton reaction (Powers et al., 2011; Bolisetty and Jaimes, 2013; Halon-Golabek et al., 2019). This is pertinent to the presence of iron within myocytes, as skeletal muscle holds the majority of the bound iron in ferritin or myoglobin, outside of the mitochondria. However, in conditions of manganese deficiency, superoxide dismutase enzymes can instead bind with copper or iron, which may restrict availability for these metals to interact with H_2O_2 , hindering its activity. When bound to these transition metals, H_2O_2 -mediated production of OH^- increases within the mitochondria, promoting

increased organelle damage and possible mitophagy (Paradies et al., 2001; Ott et al., 2007; Halon-Golabek et al., 2019).

Hydroxyl radical

During a homeostatic response, hydroxyl radicals (OH^\cdot) are the final free radicals formed as a result of ROS interaction with proteins or enzymatic degradation, and are known to promote cellular damage. Due to its electron spin configuration, OH^\cdot holds the strongest oxidizing potential of all ROS, attributing to their high reactivity. Consequently, they bind with their nearest surroundings and satisfy their lone valence electron, resulting in oxidative damage (Powers et al., 2011). Their affinity to rapidly bind to their surroundings makes it virtually impossible to directly quantify them in vitro, and only possible to evaluate products of oxidative reactions, such as disulfide bonds (Powers et al., 2011; Bekhit et al., 2013; Bolisetty and Jaimes, 2013). Due to their extreme reactivity, OH^\cdot molecules are regarded as the most damaging ROS generated in a biological setting.

Reactive nitrogen species

The term reactive nitrogen species (RNS) refers to the enzymatic development of nitrogen radicals and reactive derivatives with a nitrogen center (Powers et al., 2011; Bolisetty and Jaimes, 2013). In nature, all development of RNS is controlled by the activity of nitric oxide synthase enzymes (NOS). In mammals, NOS is generated in three distinct isoforms: neuronal (nNOS), inducible (iNOS), and endothelial (eNOS) (Kapur et al., 1997; Brannan and Decker, 2002; Förstermann and Sessa, 2012). Of these isoforms, nNOS is expressed in neurons of the central and peripheral nervous systems, and

predominantly functions for the relaxation of smooth muscle and vasodilation of blood vessels. Through immunological and cellular fractionation methods, nNOS has been detected in specialized structures of fast-twitch muscle fibers like the sarcolemma (Powers et al., 2011). Endothelial NOS is a dually acylated peripheral membrane protein found bound to caveolin-1, a caveolae structural protein associated with endothelial cells, and assists in regulating blood pressure and atherosclerosis (García-Cardena et al., 1997). Unique to muscle, immunoblotted eNOS colocalized with mitochondria of rat skeletal muscle, strongly suggesting a specific localization of eNOS to skeletal muscle mitochondria (Stamler and Meissner, 2001). Inducible NOS, while not as active in muscle, is expressed in response to stress factors such as lipopolysaccharides and cytokines (Tengan et al., 2012). When mediated by molecular oxygen, L-arginine, and electron cofactors NADPH and FAD, NOS is capable of producing nitric oxide (NO), the smallest known signaling molecule. Similar to ROS, the physiological levels of NO are recognized to have an impact on mitochondrial biogenesis, respiration, and oxidative stress. Under states of low RNS, its constituents assist in increased glucose uptake/metabolism, vasodilation, and possibly mitochondrial biogenesis (Nisoli and Carruba, 2006; Powers et al., 2011; Bolisetty and Jaimes, 2013; Hong et al., 2014). However, it is postulated that cascading of RNS can initiate nitrosative stress, producing detrimental effects on organelle function, DNA stability, and lipid/protein composition.

Nitric oxide

Nitric oxide (NO[•]) is the initial RNS formed as a result of enzymatic reactions (Figure 1). Predominantly generated using endothelial nitric oxide synthase, eNOS, NO[•]

is introduced into skeletal muscle via major arterioles surrounding muscle fibers (Bolisetty and Jaimes, 2013). Concurrently, eNOS is postulated to be found in mitochondria, sometimes identified as mtNOS. This theory is supported by the growth of enzymatic products of NO reactions (L-citrulline) in the presence of inflammatory stimuli (Stamler and Meissner, 2001). It is speculated the presence of NO^- within mitochondria is indicative of pathways independent of NOS activity such as the electron transport chain (Ghafourifar and Cadenas, 2005; Lacza et al., 2006; Arriagada et al., 2018). As a result, NO^- is capable of influencing mitochondrial composition and function. Mainly, NO^- holds a high affinity to cysteine-rich proteins, such as Complex IV-bound cytochrome c oxidase, due to the easy covalent binding, nitrosylation, of thiol groups (-SH) into disulfide bonds (Nisoli and Carruba, 2006; Tengan et al., 2012; Liu et al., 2018; Poderoso et al., 2019). Given the significance of cytochrome c oxidase and its ability to transfer multiple electrons simultaneously, accumulated NO^- present can ultimately inhibit mitochondrial respiration. Given high levels of NO^- coupled with prolonged exposure, NO can cause irreversible inhibition of mitochondrial respiration, uncoupling proteins, and ultimately resulting in apoptosis (Bekhit et al., 2013; Poderoso et al., 2019). Conversely, low levels of NO^- have been shown to trigger mitochondrial biogenesis via increased expression of Peroxisome proliferator-activated receptor gamma coactivator 1-*alpha*, PGC-1 α , encoding proliferation of mitochondrial proteins (Nisoli and Carruba, 2006; Tengan et al., 2012). Nitric oxide regulation is critical to the efficiency of mitochondrial respiration and energy production. The presence of NO^- , particularly within mitochondria, is critical considering the interaction of NO^- generates reactive derivatives of RNS. In particular, the high affinity for NO^- to bind to superoxide to

produce peroxynitrite, S-nitrosothiols (SNOs), and metal NO⁻ complexes (Stamler and Meissner, 2001).

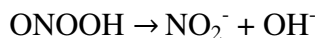
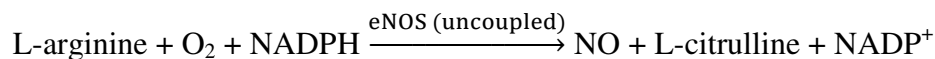


Figure 1: Generation and breakdown of reactive nitrogen species (RNS).

Peroxynitrite

During prolonged states of oxidative stress, hybridization of superoxide and nitric oxide produce peroxynitrite (ONOO⁻), a specialized molecule that can target proteins relevant to organelle stability and muscle function (Eu et al., 1999). This reaction holds a large K_m, and energetically favors production of peroxynitrite in the presence of O₂⁻ and NO⁻. As a result, the reaction is three time faster than the dismutation of superoxide to produce hydrogen peroxide, and even faster than the binding of NO⁻ with heme proteins (Powers et al., 2011). The reaction kinetics in making peroxynitrite are so favorable that accumulation of ONOO⁻ is greater than other products when O₂⁻ and NO⁻ are present. Depending on the quantity, peroxynitrite can have drastically different effects on skeletal muscle. During cellular homeostasis, Zhou et al. (2003) demonstrated peroxynitrite promotes tyrosine nitration, which in turn activates glucose uptake via lipid or protein oxidation channels by AMPK expression (Hong et al., 2014). Under conditions of reactive species amplification, ONOO⁻ supports protein nitrosylation and DNA damage (Zorov et al., 2000; Stamler and Meissner, 2001; Bekhit et al., 2013) Specifically, ONOO⁻ targets proteins rich in cysteine and methionine residues, as these amino acids hold high oxidative potential due to the weak stability of their thiol groups (-SH). These

reactions result in protonated peroxynitrite formation (ONOOH), facilitating an energetically favorable decomposition of peroxynitrite to produce hydroxyl radicals ($\text{ONOOH} \rightarrow \text{NO}_2^- + \text{OH}^\cdot$) (Wang et al., 2002). As a result, peroxynitrite is not only highly disruptive towards protein function, but an effective precursor to the development of highly reactive oxygen species.

Hyperchlorite

Hyperchlorite is a myeloperoxidase formed by binding hydrogen peroxide with molecular chloride (Cl^-). Similar to peroxynitrite, hyperchlorite is capable of damaging biomolecules via oxidizing lipids, electron donor activity, and thiols. In rare cases, hyperchlorite can be converted to hypochlorous acid, which can diffuse across cellular membranes and is acidic enough to cause further protein and lipids damage (Powers et al., 2011). However, this RNS is found in infection-responding neutrophils, specialized white blood cells, and is commonly seen when experimentally-induced stress in bacterial species. Due to the irregular generation and unknown effect in mammalian cells compared to well-known RNS, it is regarded as a less critical RNS in the respect of skeletal muscle damage.

Redox-sensitive signaling pathways

While it is still unknown how to define distinct levels of ROS based on their biological effects, there is evidence that ROS/RNS trigger downstream signaling pathways that stimulate cell growth, differentiation, proliferation, and apoptosis. These pathways can also act as precursors to generate enzymes and antioxidant factors which

can suppress the quantity and severity of reactive species (Miller et al., 1993). Often, these pathways are triggered by the mere presence of ROS/RNS generated, and are sometimes expressed due to the change in activity of distinct proteins (cysteine, kinases, phosphatases) linked to oxidation or nitrosylation products (Bolisetty and Jaimes, 2013). Concurrently, there is evidence to suggest that the *degree* of ROS/RNS damage may alter the level of expression across redox-signaling pathways. Simply put, the expression of redox-signaling pathways is not positively correlated [linearly related] with the sheer amount of ROS/RNS generated (Bolisetty and Jaimes, 2013). Given the diversity of pathways activated in response to ROS/RNS, it is challenging to understand the intricate relationship between reactive species and biological response pathways. As a result, it is pertinent to understand the known connection between these foundational pathways and oxidative stress factors.

Nuclear factor erythroid 2-related factor (Nrf2)

During ideal physiological conditions, the production of ROS generated by oxidative phosphorylation is easily suppressed. This is predominantly due to the activity of the nuclear factor erythroid 2-related factor (Nrf2), which is responsible for the regulation of antioxidants and radical scavengers produced *in vivo* (Bolisetty and Jaimes, 2013; Kozakowska et al., 2015). Stored in the cytosol, Nrf2 is sequestered to Kelch like-ECH-associated protein 1 (KEAP1). A sensor protein, KEAP1 is designed to inactivate in response to reactive oxygen species (Dodson et al., 2015). When present, H₂O₂ and NO have been shown to inactivate KEAP1 by oxidizing several cysteine residues present, stimulating Nrf2 to enter the nucleus and bind to the antioxidant response element (ARE)

attached to stress responsive genes. This generates transcription factors needed to signal release of antioxidants (Fourquet et al., 2010; Bolisetty and Jaimes, 2013). It is possible that Nrf2 activators may actually increase ROS/RNS generation (Fourquet et al., 2010), however, the evidence is inconclusive. While Nrf2 activity is a critical mediator to antioxidant signaling during low levels of reactive species, its activity is generally disregarded compared to other signaling pathways during increased cellular stress (Fourquet et al., 2010; Bolisetty and Jaimes, 2013).

NFκB/AP-1

As expected, a multitude of physiological pathways work dependently, as the activities of certain proteins for one pathway are mitigated by the expression of transcription factors of another. For example, the function of Activator Protein 1 (AP-1) is largely dependent on the signaling pathways of NFκB (nuclear factor kappa-light-chain-enhancer of activated B cells) and MAPK (mitogen-activated protein kinase). In both the cytoplasm and nucleus, NFκB and AP-1 react to “moderate” amounts of ROS. Concomitantly, each are stimulated by the presence of H₂O₂ and a variety of cytokines related to inflammation and apoptosis, such as Interleukin-1/6, and Tumor necrosis factor- α (Powers et al., 2011). In concert, both pathways bind to DNA promoters to activate antioxidant proteins including superoxide dismutase-2 (SOD2), cyclooxygenase-2, catalase (CAT), MnSOD, and glutathione peroxidase (GPx) (Bolisetty and Jaimes, 2013). Interestingly, NFκB also assists in activating iNOS, which suggests that it indirectly promotes expression of other redox-signaling pathways by the generation of NO (Nisoli and Carruba, 2006; Powers et al., 2011). While the activation of this pathway

is shown to develop antioxidant combatants against reactive species, NF κ B is implicated to increase inflammation via upregulation of enzymes that facilitate ROS production such as NADPH oxidase and xanthine oxidase, as well as NOS isoforms (Bolisetty and Jaimes, 2013). As a result, an intricate balance of NF κ B and AP-1 regulation is necessary as too little expression will not suppress reactive species and over expression may generate high enough levels of ROS to oxidize proteins relevant to cellular structure and organelle function, reducing their activity entirely.

Mitogen activated protein kinase (MAPK)

From another perspective, MAPK promotes cell proliferation and differentiation in response to cellular damage as a result of reactive species (Dröge, 2002). In regards to cell turnover, MAPK pathways activate dormant satellite cells, the precursor to active skeletal muscle cells (Jones et al., 2005). This is accomplished by the diverse composition of MAPK phosphate subsets, each can be grouped based on their subcellular localization (i.e., MAPK-2 found in the nucleus; MAPK-3 present in the cytoplasm). These subsets act to dephosphorylate MAPK, inactivating it. Interestingly, ROS can target MAPK phosphates during states of high ROS production, and modify their function to upregulate MAPK activity (Son et al., 2011). Upregulation of MAPK is speculated to initiate translocation of heat shock proteins 25/27 to the myofilament Z-disk to prevent apoptosis (Blunt et al., 2007). In turn, there is a tight and intricate relationship between MAPK activation and ROS production.

Homeostatic response

In order to maintain cellular homeostasis, muscle cells have evolved to use a network of different antioxidant defense systems to counteract oxidative damage in the presence of ROS. The diverse antioxidant systems distribute enzymes, which are specific to certain reactive species and designed to produce a subsequent reactive specie which is specific to another antioxidant. This cascade of enzymatic reactions intend to produce final products that are stable for ideal cellular function, such as water or molecular oxygen. Intrinsic regulation of ROS includes both enzymatic and non-enzymatic antioxidants and these regulatory systems are located across both intracellular (organelles; cytoplasm) and extracellular space. The regulatory mechanism for ROS depends on the antioxidant. Predominantly, endogenous antioxidant systems convert ROS to less reactive states, but can also prevent ROS propagation by minimizing available pro-oxidants such as chelating iron and copper (Miller et al., 1993). Interestingly, antioxidant activity varies across different muscle fibers, with oxidative (Type I) fibers noticeably greater than glycolytic (TypeII) fibers (Powers et al., 1994). This is understandable as Type I fiber types have more mitochondria. Dietary antioxidants (α -tocopherol) can act as ROS scavengers and eliminate ROS molecules (Suman et al., 2014). Antioxidants are readily present endogenously and mitigate cellular response to ROS. In similar fashion, exogenous enzymes can be administered to combat ROS, as Batifoulier et al. (2002) showed greater membrane stability of organelles as supplemented α -tocopherol inclusion rates increased. Typically, such antioxidants are provided in daily rations, as there is potential that supplementation can also hold benefits of energy efficiency and growth in livestock (Miller et al., 1993).

Superoxide dismutase

As the name implies, endogenous antioxidant enzymes are those which are produced within an organism to provide protection against lipid, protein, or nucleic acid oxidation. In relation to reactive species, superoxide dismutase (SOD) is the antioxidant that starts the cascade of enzymatic reactions to eliminate ROS. Discovered in 1969, SOD dismutates O_2^- radicals to form H_2O_2 as seen in Figure 2 (Paradies et al., 2001; Descalzo and Sancho, 2008; Delliaux et al., 2009). Superoxide dismutase comprises three isoforms, all of which use unique cofactors to facilitate the breakdown superoxide anions. Superoxide dismutase-1 is located in the cytosol and inner membrane space, and uses a copper-zinc binding cofactor. Superoxide dismutase-3 uses the same cofactor but is found in the extracellular space outside a muscle cell. Superoxide dismutase-2 uses a manganese cofactor and resides in the mitochondrial matrix, making it the primary isoform used to prevent ROS accumulation during oxidative phosphorylation (Miller et al., 1993). When examining skeletal muscle, the general activity of SOD varies by location, with ~15-35% of SOD activity occurring within the mitochondria and the remaining activity in the cytosol. This implies that if ROS generation occurs outside of the mitochondria, there is sufficient SOD that resides outside of the mitochondria to break down O_2^- . With sufficient supply of SOD, prevention of ROS formation is attainable, eliminating the onset of oxidative stress from the start.

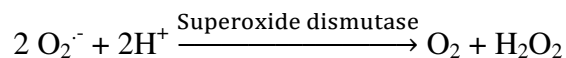


Figure 2: Superoxide breakdown and production of hydrogen peroxide.

Glutathione peroxidase/catalase

Upon completion of SOD-mediated degradation, ROS species are further broken down to water molecules via glutathione peroxidase and catalase (GPx and CAT, respectively; Figure 3) (Gatellier et al., 2004). When examining the two enzymes, GPx appears to be relatively similar to SOD in respect to its diverse isoforms and their localization in muscle cells. In particular, GPx activity can greatly influence the mitochondrial permeable transition pore due to the large quantity of cysteine residues present (Dalle-Donne et al., 2003). Additionally, GPx requires both a selenium cofactor and glutathione (GSH), a cysteine based protein, which supplies electrons needed to trigger a forward reaction. This may act as a detriment to GPx activity however, as the cell must have a redox cycle to have the oxidized glutathione (GSSG) to return to its reduced state (Descalzo and Sancho, 2008). Catalase, like SOD, uses iron as a required cofactor to reduce H_2O_2 to water and molecular oxygen (Zámocký and Koller, 1999; Kirkman and Gaetani, 2007). Compared to GPx, CAT only uses one cofactor, iron, to regulate its activity instead of a reduced, thiol-based peptide. Conversely, CAT efficacy is more variable, as its concentration is positively correlated to its affinity towards H_2O_2 degradation (Pradhan et al., 2000). This implies that CAT is most effective against H_2O_2 once the ROS has grown in quantity rather than when it becomes available for enzymatic degradation. While both enzymes act to degrade the same reactive species, both GPx and CAT are ineffective in certain conditions. However, its counterpart remains active in that same situation. This results in a relationship between each enzyme, where conditions which may inhibit optimal activity of one are more conducive for the activity of the other, which helps maximize suppression of ROS-mediated damage to myocytes.

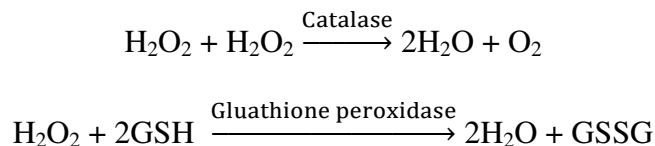


Figure 3: Enzymatic breakdown of hydrogen peroxide using catalase and glutathione peroxidase.

Alpha-tocopherol (vitamin E)

Dietary supplementation of vitamin E is widely regarded as the best applicable method to delay meat discoloration and lipid oxidation (Faustman et al., 1998). When consumed, α -tocopherol is predominantly deposited in the membranes of major organelles. Derived from the carotenoid family, α -tocopherol is a lipid-soluble antioxidant consisting of long chains of conjugated double bonds and a benzene ring, and acts as an effective inhibitor of free radical-induced peroxidation of unsaturated fatty acids (Suman et al., 2014). This prevention occurs by terminating free radical reactions during the propagation of lipid radicals, oxidizing α -tocopherol into a tocopherol radical (Ryan et al., 2010). Redox reactions can occur using ascorbate and reduced glutathione to recycle this radical back to α -tocopherol, allowing further ROS scavenging (Packer et al., 1979; Ryan et al., 2010). Given the capacity for carotenoids to be deposited within organelle membranes responsible for myofibrillar function, their presence can profoundly affect protein and lipid oxidation (Harris et al., 2001).

Mechanism of oxidative stress

Aerobic organisms generate ROS as a result of oxidative phosphorylation. As a safeguard, generation of a multitude of antioxidants and enzymatic compounds are utilized to suppress and eventually remove any detrimental effect of these ROS.

However, evidence has shown instances where physiological response mechanisms are not sufficient in combating ROS, such as inflammation, chronic disease, illness, or muscle injury. As a result, these occurrences can stem from the phenomena known as oxidative stress (Figure 4). As its name implies, oxidative stress is defined as the imbalance between the production of ROS and antioxidant defenses, in which oxidative stress favors ROS (Ott et al., 2007; Tsutsui et al., 2011). The exhaustion of defense mechanisms allows ROS to react with their surroundings, along with each other, resulting in damage to proteins, lipids, and DNA. With sustained damage to cellular structures and altered redox-reactions, oxidative stress worsens (Miller et al., 1993; Ott et al., 2007; Celi, 2010; Scicchitano et al., 2018). Under states of prolonged oxidative stress, a timeline which is undefined, oxidative stress changes cellular function and the biological mechanisms used to sustain cellular development, validating the need for further investigation into oxidative stress effects.

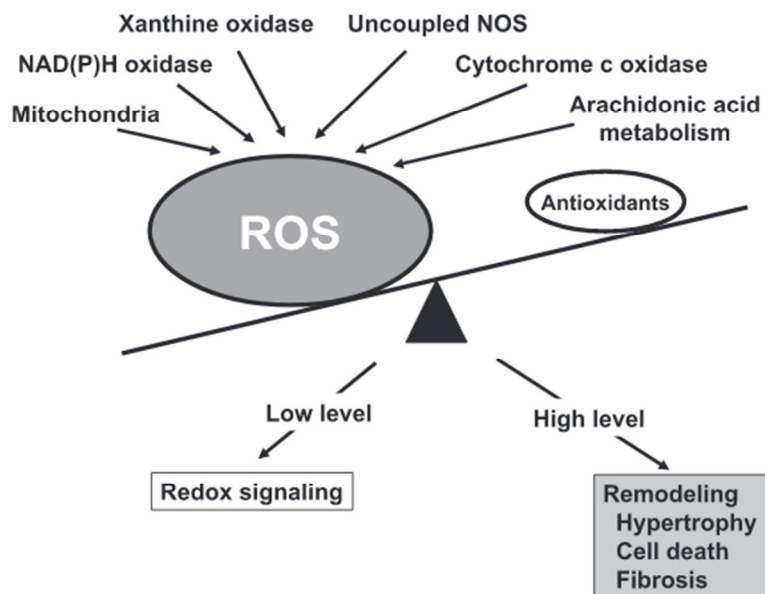


Figure 4: Enzymatic sources of reactive oxygen species (ROS) and their pathophysiological role involved in oxidative stress (Tsutsui et al., 2011).

DNA profile

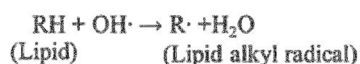
Various components of DNA can be modified via oxidative stress. In particular, the oxidation of purine and pyrimidine bases making up the foundation for the deoxyribose backbone are the most susceptible to oxidative alteration (Bekhit et al., 2013). This promotes permanent cross-bridging of these nucleic acids,, drastically changing the functionality of the DNA molecule, forcing recycling of these molecules and slowing cellular nuclei to meet cellular demands for DNA turnover. In particular, the damage by ROS to mitochondrial DNA disrupts transfer RNA and ribosomal RNA, impacting protein synthesis and essential ATP production by the electron transport train. Oxidative damage induced by ROS is likely a major source of genomic instability leading to respiratory dysfunction (Ott et al., 2007). Oxidative stress-mediated DNA damage is more commonly linked to chronic illness and has not yet been linked to meat quality. A more in-depth investigation into the presence of oxidative stress and its impact on the formation of proteins using transcriptomic profiling is necessary.

Lipid oxidation

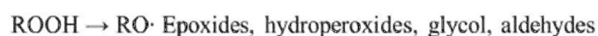
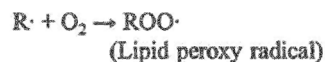
Oxidation of lipids is primarily focused on polyunsaturated fatty acids (PUFAs). Due to weak allylic central carbon binding in double bonds, PUFAs contain methylene bridges, which hold highly reactive hydrogen atoms. In the presence of ROS, the lipid oxidation mechanism can begin (Figure 5). Known as the initiation step, fatty acids lose a hydrogen ion during oxidation to produce highly unstable lipid radicals called hydroperoxides (Bekhit et al., 2013; Domínguez et al., 2019). In regards to fatty acids, PUFAs are the most susceptible to lipid oxidation. In the presence of molecular oxygen,

hydroperoxides react to produce lipid peroxy radicals. Due to the instability of these radicals, they behave like ROS to scavenge an available hydrogen from their surroundings. Ironically, their surroundings are other fatty acid structures. This cascade of lipid radical binding to non-radical fatty acids begins an exponential propagation in the oxidation of surrounding lipids. Eventually, a localized area of fatty acids is overwhelmingly comprised of lipid radicals, allowing these radicals to bind with one another, producing lipid peroxides (Domínguez et al., 2019). Concurrently, the generation of lipid oxidation promotes further biochemical changes, such as increased membrane fluidity, physiological function, enzyme inactivation, and protein denaturation (Stark, 2005; Ott et al., 2007; Powers et al., 2011; Bekhit et al., 2013; Kozakowska et al., 2015; Scicchitano et al., 2018), all of which, can result in major shifts in cellular vitality and stability.

Initiation step



Propagation step



Termination step

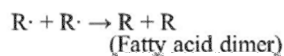


Figure 5: Systematic process of lipid peroxidation.

Lipid oxidation interrelationship with meat color

Fresh meat color is one of the largest indicators of freshness to consumers and the primary factor in consumer purchasing (Gatellier et al., 2001). Consumer analysis indicates that a 20% discoloration of meat during retail display can reduce total beef sales by 50% (Hood and Riordan, 1973). The color of meat has been related to the combination of three states of myoglobin on the surface of meat: reduced myoglobin, oxymyoglobin, and metmyoglobin. The overall composition of myoglobin pigments depend on the rates of oxymyoglobin oxidation and metmyoglobin reduction (Gatellier et al., 2001), which has been linked to lipid oxidation (Faustman et al., 2010). Intermediate radicals are produced as a result of oxidative propagation of lipids and can accelerate myoglobin oxidation, increasing meat discoloration. Wong (1989) explained how iron-bound myoglobin can catalyze lipid oxidation, indicating that ferric (Fe^{3+}) can promote lipid autoxidation, generating superoxide anions (O_2^-). When reacting with protein-bound thiol groups, additional O_2^- generation can occur. Concurrently, O_2^- acts as a byproduct of redox reactions, rapidly being dismutated to H_2O_2 , which can further increase lipid and myoglobin oxidation simultaneously.

Protein oxidation

Recent literature has been recognizing the emerging investigation in the oxidation of proteins sources in food. The growing interest stemmed from the idea of protein sources to be oxidized via ROS reactions, with ROS potentially causing deleterious effects on muscle proteins alongside lipids (Stadtman and Levine, 2003). In turn, the

investigation into how protein oxidation occurs, and how oxidative stress can promote these reactions, has yet to be adequately defined.

Using current knowledge, protein oxidation can occur through the targeting of peptide backbones and specific amino acid side chains, resulting in a loss of sulfhydryl groups (protein cross-linking) and generation of carbonyl derivatives (Estévez, 2011). In general, protein oxidation occurs similar to oxidation of lipids. Using the generation of ROS, oxidative radicals target the functional groups along the side chain of amino acid residues and scission of peptide backbones. In regards to lipid and protein oxidation, radical formation can be a result of the Fenton reaction, the catalysis of transition metals iron/copper in the presence of H_2O_2 radicals (Martinaud et al., 1997; Estévez, 2011). When oxidative radicals are reduced, the transfer of energy with the hydrogen atoms can generate protein radicals. Protein radicals further react with each other and molecular oxygen to form a population of radicals and generate hydroxyl derivatives of proteins (Stadtman and Levine, 2003). The routes taken to generate these protein derivatives are dependent on the specific amino acid complexes which are targets by ROS, and thus, each method of protein oxidation can merit different products from protein oxidation.

From protein oxidation, there are a number of different derivatives which can be used to evaluate oxidation. Of these, loss of sulfhydryl groups (R-SH) is a common measure of protein oxidation. While most thiols are low-reacting given their pK_a (~8.5), certain amino acids can create a charged environment for thiolate anions to exist. These anions are more readily oxidized/nitrosylated, especially in the presence of ROS/RNS (Dalle-Donne et al., 2003; Ying et al., 2007). As a result, cysteine and methionine are easily oxidized than others due to their thiolate side chains. If oxidized, thiol groups tend

to bind with their cysteine-oxidized thiol counterparts, forming covalent disulfide cross-bridges (Frederiksen et al., 2008; Estévez, 2011). These cross-bridges are more stable than most cross-linkages in amino acids, due to their strong attraction from sharing electron orbital with one another, and require exogenous chemical conditions to prevent their formation (i.e., beta-mercaptoethanol). During conditions which promote protein denaturation (cooking, high pH, or oxidative stress), disulfide bonds are very common and act to change the conformation of protein shape, and possibly function (Soladoye et al., 2015). As a result, the change in composition drastically alters protein stability, acts as a severe detriment to protein function and quality, and can affect downstream quality attributes of muscle postmortem.

In addition to thiol oxidation, generation of carbonyl derivatives is a well-known indicator of protein oxidation. Carbonylation is the irreversible, non-enzymatic formation of aldehydes and ketones during protein oxidation (Estévez, 2011). Carbonyl derivatives are commonly formed from amino acids lysine, threonine, and arginine using metal-catalyzed oxidation against amine groups (Stadtman and Levine, 2003). After oxidation, amino acids convert into irreversible aldehyde groups. Given the moiety of carbonyls, they are capable to continue binding with surrounding peptide bonds to form cross-links via Schiff-base formation, which increases the susceptibility of neighboring peptide to form ketones or aldehydes. Unique to carbonylation, myoglobin is susceptible to protein oxidation when exposed to H_2O_2 due to its binding with transition-metals iron or copper. As a result, oxidation reactions occur to form metmyoglobin pigments. Estévez and Heinonen. (2010) discovered H_2O_2 -mediated metmyoglobin formation to produce

hypervalent radical ferrylmyoglobin, which has been linked to lipid and protein oxidation (Baron and Andersen, 2002).

Stability of the sarcoplasmic reticulum

As previously stated, the sarcoplasmic reticulum (SR) acts as a regulatory organelle for muscle contraction by the release and sequestering of calcium ions into the cytosol by the ryanodine calcium channel and SERCA, respectively. (Rossi and Dirksen, 2006). Consequently, the SR acts as a critical component to meat quality, as calcium acts as a catalyst for the activation of proteolytic calpains used in meat tenderization (Koochmaraie, 1996). Oxidation of the SR membrane could increase calcium flow out of the SR. This effect occurs as a result of prolonged postmortem muscle aging in conjunction with chronic illness, when membrane repair mechanisms are minimal (Fulle et al., 2004). Certain ROS can influence calcium homeostasis across the sarcoplasmic reticulum and mitochondria. Isaeva et al. (2005) reported that H₂O₂ increased calcium release from the sarcoplasmic reticulum. Other factors, such as prolonged exercise, have been shown to produce ROS (Li et al., 1999).

During states of oxidative stress, nitric oxide is generated and reacts with protein residues found within the SERCA and ryanodine receptors. Cysteine and methionine residues are largely comprised of oxidatized sulfhydryl-groups. When oxidized, SERCA activity is inhibited via disulfide formation, drastically decreasing uptake of cytosolic Ca²⁺ ions and consequent inhibition of muscle contraction (Powers et al., 2011). Ishii et al. (1998) revealed the reversible activity of SERCA inhibition after the loss of nitric

oxide. However, this recovery of SERCA activity is quite slow considering the reactivity of NO occurs within seconds.

Activity of ryanodine receptors is likely dependent on *S-nitrosylation* of amino acids with high oxidative potential. Low concentrations of NO-mediated oxidation did not affect RyR1 channel activity, as the binding of radical hybrid peroxynitrite forms permanent cross-links of disulfide bonds within RyR proteins. Interestingly, subunit cross-linking did not inhibit activation of skeletal RyR channels by disulfide bond formation from sulfhydryl groups. Simply put, low NO concentration didn't stop activity of ryanodine receptors, but did alter the functionality of receptors, changing their function and preventing physical release of Ca²⁺ ions. Conversely, high concentrations of NO could activate RyR channels. At high enough concentrations, diversity of NO-related species are created after binding to RyR (GSNO, CysNO, and SIN-1) and their concentration ratio regulated the denitrosylation of RyR (Ishii et al., 1998; Eu et al., 1999). It is not quite understood how the diversity of these NO-related species are controlled, as it is likely dependent on factors such as cellular composition, organelle membrane structure, NO concentration, and antioxidant interaction (Eu et al., 1999).

Mitochondria stability

Multiple facets of cellular life and death revolve around mitochondrial activity. Not only are mitochondria the center for ATP production, they are also the center of intracellular ROS production (Bolisetty and Jaimes, 2013). During bouts of oxidative stress, the integrity of mitochondrial function is jeopardized both within the matrix and across the extracellular membrane. Ott et al. (2007) demonstrated how aconitase, an iron-

sulfur bound protein, acts as a catalytic converter of citrate to isocitrate in the Krebs cycle. Irreversible oxidation of aconitase during oxidative stress caused dysfunction of the Krebs cycle and diminished ATP production due to the lack of electron donors generated. This progressively decreased the membrane potential of the mitochondrial matrix, and if not resolved, caused the degradation of the matrix and loss of mitochondrial function to promote apoptosis (Zorov et al., 2000; Sierra and Olivan, 2013). Free fatty acid production as a result of membrane oxidation confounds the loss of membrane potential. Bernardi et al. (2006) stated the combination of increased ROS production and lowered membrane potential activated the MPTP in a last-ditch effort to expel ROS and prevent mitochondrial destruction. However, the increased proportion of ROS produced, combined with the depolarization of mitochondria membrane, facilitated slow efflux of ROS, which can result in mitophagy (Di Lisa et al., 2001). These cascading events all promote cellular death and initiate apoptotic pathways, which are responsible for replacing oxidized proteins and organelles.

Apoptotic pathway

Mitochondrial ROS generation, coupled with other pro-apoptotic proteins, are the central cause of apoptosis, or the execution of programmed cellular death. The apoptotic process is a non-inflammatory pathway complementary to mitosis, which acts in concert with enzymatic proteins. The end goal is to break down necrotic cells and induce growth of new cells (Ott et al., 2007). At mitochondrial failure, key proteins are utilized in cell signaling of apoptosis and can be evaluated as markers of apoptotic pathways during stages of oxidative stress (Dröge, 2002).

Cytochrome c

In living cells, cytochrome c functions as a the primary electron carrier between Complex III and IV of the electron transport chain (Ke et al., 2017). As it is related to the electron transport chain, cytochrome c is localized within the inner mitochondrial membrane and bound alongside cardiolipin, an anionic phospholipid that assists in stability of the mitochondrial membrane. Ott et al. (2007) postulated cardiolipin and cytochrome c were so tightly bound that the separation requires major changes in cardiolipin's affinity for cytochrome c to be released. In states of oxidative stress, the oxidation of cardiolipin facilitates cytochrome c release and is shown to assist in the expulsion of cytochrome c from the inner mitochondrial membrane (Kagan et al., 2009). Concomitantly, pro-apoptotic proteins (Bax) allow the opening of mitochondrial pores along the outer membrane, allowing cytochrome c release from the mitochondria into the cytoplasm. Once in the cytoplasm, cytochrome c initiates gene signaling of proteolytic proteins used to complete apoptosis.

Caspase system

Later stages of apoptosis facilitate the degradation of necrotic cells in order to provide growth for new and adapted structures. Madesh and Hajnóczky (2001) commented on how the presence of cytochrome c in the cytoplasm signaled the family Apaf-1 genes (apoptotic protease activating factor 1), which activate enzymatic proteins. Correspondingly, Apaf-1 binds with cytosolic cytochrome C to form the apoptosome, a large quaternary protein that enables the activation of caspase proteins (Madesh and Hajnóczky, 2001). The caspase work in conjunction to achieve proteolytic function.

Sierra and Olivan (2013) reviewed distinct groups of caspases and their role in apoptotic pathways. To start, initiator caspases 8, 9, and 12 turn on in response to apoptotic stimuli and accordingly activate effector caspases 3, 6, or 7. This is completed by the progressive cleaving of each type of caspase in order to cleave subsequent caspases, activating each upon being cleaved. When activated, the caspase enzymes initially function to cleave Poly (ADP Ribose) Polymerase, designed to repair proteins, allowing natural cellular breakdown and eventual death to be met (Porter and Jänicke, 1999). Additionally, caspase has been theorized to cleave calpastatin, an inhibitor of postmortem aging in muscle. Wang et al. (1998) showed calpastatin activity held a negative relationship with caspase activity up to two days postmortem at refrigerated temperatures. Caspase degradation of organelle membranes and cytoskeletal proteins increases cytosolic calcium levels and calpain activity (Wang, 2000). This action assists in early postmortem tenderization.

Small heat shock proteins

Inherently, biological homeostasis of structural proteins is maintained through the use of chaperone proteins. As its name suggests, chaperone proteins stabilize unstable proteins and oversee the correct assembly of protein growth (Creagh et al., 2000; Lomiwes et al., 2014). In relation to oxidative stress, the primary isoforms of interest are small heat shock proteins (sHSPs). Of sHSPs, HSP20 (20 kDa), HSP27 (27 kDa) and $\alpha\beta$ -crystallin (22 kDa) have been implicated in meat quality related to oxidative stress. Fischer et al. (2002) found that the expression of these chaperone proteins in skeletal muscle occur at a high volume compared to other tissues. Due to the frequent generation

of oxidative stress factors in skeletal muscle, it makes sense that increased sHSPs are produced as a response to apoptosis. During homeostatic conditions, sHSPs can act as regulatory factors in myofiber composition. Lomiwes et al. (2014) stated that sHSPs regulate the growth of actin myofilaments. During the conversion of G-actin to F-actin, the positive terminal associated with F-actin growth has an affinity ten times greater to bind G-actin than the negative end. As HSP27 inherently binds to the positive end, inhibiting exponential growth, phosphorylation of HSP27 is required to be released to allow for further F-actin growth. This action prevents the overproduction of actin filaments. Der Perng et al. (1999) illustrated how HSP27 works alongside $\alpha\beta$ -crystallin to prevent overgrowth of intermediate desmin filaments, controlling their growth to replace damaged proteins. During oxidative stress, however, they function as modulators of the apoptotic mechanism. Creagh et al. (2000) stated an increase in stress factors (ROS, heat) promote heat shock factors (HSF) to transport into the nucleus, bind with HSP genes, and upregulate HSPs to combat apoptotic mechanisms. In particular, HSP27 can bind to cytosolic cytochrome C and other pro-apoptotic proteins, inhibiting apoptosome formation and subsequent activation of caspase enzymes. However, the overexpression of these sHSPs can prevent myogenic differentiation necessary for new cellular growth, as shown by $\alpha\beta$ -crystallin inhibition of caspase-3 dimers to prevent degradation of necrotic cells. This reinforces the rule of sHSPs as a necessity during oxidative stress response, regular cellular growth, and development.

Enzymatic aging

Cellular apoptosis begins at death, only there is no longer a supply of oxygen present to sustain energy production for the repair, growth, and protection for cellular turnover. Consequently, it has been suggested that apoptotic factors may contribute to initial postmortem degradation of proteins during the conversion of muscle to meat (Ouali et al., 2006). Cao et al. (2010) found the generation of caspase-9 activity increases exponentially from 4 to 30 hours postmortem, and was shown to peak in its activity for as long as 7 days postmortem. Total activity of caspase activity may also be muscle-specific, as different muscles are comprised different fiber types. As the degree of apoptosis is correlated to mitochondrial activity, different fiber types require varying number of mitochondria for normal muscle function *in vivo*. Therefore, fiber type may affect mitochondrial-mediated apoptotic processes in the early postmortem development of caspase (Kemp and Parr, 2012). Early increase in apoptotic activity is logical, as death coupled with anaerobic conditions and the lack of ATP generation would increase caspase activity, and any external injury or advanced oxidative stress could further impact the extent of apoptosis early postmortem. Huang et al. (2016) reported that increased postmortem aging of beef skeletal muscle increased the ratio of cytosol:mitochondria cytochrome c and decreased anti-apoptotic factor HSP27, indicating continual proteolytic degradation as aging continues. Laville et al. (2009) reported that samples from bulls exhibiting less shear force (tender; 27.7 N) compared to samples with more shear force (tough; 41.2 N) 7 days postmortem, showed a greater amount of inner and extracellular mitochondria fragments in samples collected 10 minutes postmortem. Thus, increased aging results in faster degradation of organelle

structures. With the help of increased organelle degradation, downstream mechanisms contributing to meat tenderness may become more active early postmortem.

Calpain-calpastatin relationship

At the onset of postmortem tenderization, changes in muscle pH, temperature, and enzymatic inhibitors all contribute to the behavior of proteolytic enzymes to progressively degrade muscle structure as a natural degradation process. Under controlled conditions, this is a vital mechanism to enhance the palatability of fresh meat (Koochmaraie and Geesink, 2006). Postmortem improvement in meat tenderness results from the breakdown of myofibrillar structures by endogenous enzymes (Ouali et al., 2006). The primary enzymes recognized to contribute to postmortem aging are calpains, cathepsins, and, to an extent, 20S proteasome. Of these, calpains are by far the most extensively researched family of enzymes linked to meat tenderness. While Pandurangan and Hwang (2012) comment on the ability of calpains to modify proteins *in vivo*, their primary effect on meat quality is the overarching mechanism responsible for tenderization in muscle proteins (Kemp and Parr, 2012; Lian et al., 2013). Calcium-dependent, non-lysosomal peptides, the three subsets of calpains within skeletal muscle are μ -calpain, m-calpain, and calpain 3. In regards to skeletal muscle postmortem, μ -calpain and m-calpain take precedent, as calpain 3 is not associated with calpain inhibitors and functions for maintenance of the structural protein titin (Ilian et al., 2004). When active, the calpain system targets substrates that relate to structural integrity of the sarcomere. In particular, the hydrolysis of desmin, titin, and nebulin are recognized as the first proteins degraded as a result of postmortem aging (Kemp et al., 2010; Lian et al.,

2013; Bhat et al., 2018). These proteins assist in the sarcomeric alignment of the myofibril during contraction, and maintain the binding of the Z-line to both myofibrils and costameres connected to the sarcolemma (Huff Lonergan et al., 2010). The hydrolysis of these proteins results in the detachment of sarcomere-sarcolemma binding in addition to sarcomere-sarcomere adhesion found at the Z-disk (Taylor et al., 1995; Henderson et al., 2017) Of the two active isoforms, μ -calpain is the more readily active due to its lower molar calcium concentration needed to function ($\sim 5\text{-}50\mu\text{M}$) compared to m-calpain ($\sim 200\text{-}800\mu\text{M}$). Taylor et al. (1995) reported μ -calpain to be more active in early postmortem compared to m-calpain, when the majority of postmortem proteolysis of key myofibrillar proteins including nebulin, titin, desmin and troponin-T occurs. Their activity is short lived, however, as their unstable state, combined with calpain inhibitor calpastatin, quickly suppress the majority of their activity within a few days postmortem. Conveniently, the progression of postmortem aging results in greater calcium leakage from the sarcoplasmic reticulum and mitochondria, the same organelles responsible for muscle contraction, facilitating m-calpain activity.

In addition to cytosolic calcium levels, the activity of calpain is regulated by calpastatin (Ouali et al., 2006). Calpastatin (110 kD) is the inhibitor specific for μ - and m-calpain and coexists with calpain. Calpastatin is comprised of four domains on its N-terminal, each of which can independently inhibit calpain, meaning a single calpastatin is capable of inhibiting four calpain units (Ouali et al., 2006; Pandurangan and Hwang, 2012). In relation to oxidative stress in terminal patients, calpastatin is shown to prevent calpain activity during apoptosis, inhibiting the dysregulation of harmful cells and promoting chronic inflammation (Momeni, 2011). In the same manner, apoptotic

pathways resulting in cellular degradation upregulate calpastatin in response to calpain activity. Ouali et al. (2006) and Kemp and Parr (2012) commented on the proportional growth of both calpain and calpastatin to be fairly consistent regardless of muscle or fiber type, insisting that calpain tenderization can be limited because of calpastatin inhibition regardless of length of postmortem aging. The literature has shown an overall decline in change of shear force in meat as aging progresses. Given the minimal change of shear force after a certain time, prolonged aging could promote more negative effects to meat quality, such as shorter shelf life and greater discoloration. Concurrently, prolonged aging will eventually create no significant difference among consumer tenderness evaluation (Miller et al., 2001; Perry et al., 2001). It is recognized, therefore, that early postmortem aging of muscle holds the greatest impact on total tenderization in lowering shear force of meat.

Cathepsins

First discovered in 1950, cathepsins were the first enzymatic system postulated as the postmortem tenderization mechanism (De Duve et al., 1955; Ouali et al., 2006). Cathepsins are a family of enzymes stored within the lysosome, and are identified based on the frequency of amino acid residues within each type, such as cysteine, aspartic, and serine groups. In particular, cathepsin L and B, both with the cysteine active site, are implicated to increase tenderization by hydrolysis of desmin, troponin, nebulin, titin, and tropomyosin. However, extensive research correlating cathepsins to meat tenderness is inconclusive (Taylor et al., 1995). Koohmaraie et al. (1991) showed that there is minimal degradation as a result of cathepsins. This is because tenderization is not associated with

actin and myosin degradation, both of which are primary substrates for cathepsins (Okitani et al., 1980). This is coupled with the multitude of hurdles that must be met in order to allow cathepsin activation, including breakdown of the lysosome. Koohmaraie (1988) showed cathepsins to have an optimum activity in a more acidic (pH ~5.0) environment. Zamora et al. (2005) suggested the inhibition of cathepsins, via cystatin, to be the superior indicator of meat tenderness, as there is a stronger correlation between activity of cysteine cathepsin inhibitors and meat toughness than there is cathepsin enzymes and meat tenderness. However, this correlation was made with a 6-variable regression equation comprised of factors such as rate of pH decline, decline of μ -calpain level, and lactate dehydrogenase activity.

Manipulation of oxidative stress precursors

Generation of oxidative stress factors are contingent on the onset of physiological stress conditions. While there is inherent generation of ROS due to oxidative phosphorylation, the generation of ROS can result from mitochondrial processes influenced by extrinsic factors. While their degree of influence on oxidative stress is uncertain, it is well-recognized that external factors can be integral to oxidative mechanisms related to meat quality, most notably protein oxidation and lipid peroxidation.

Diet

In animal production, diet composition has always been a topic of discussion. Due to the variety of animal diets available, diet composition is commonly developed due to

cost, nutritional value, and known health and performance benefits that can benefit the growth and development of livestock. There is an ever-present need to understand how commercial diets may influence animal performance, feed efficiency, and even meat quality.

In commercial agriculture, the majority of intensive finishing systems in the United States incorporate energy-dense concentrates such as grains supplemented with essential minerals and fibers. This strategy is known to assist in increased weight gain and muscle growth in livestock compared to finishing strategies focused on grass-fed diets. In turn, diets impact fatty acid composition of muscle cells and organelle structures. Hwang and Joo (2017) demonstrated how cattle from different genetic and environmental backgrounds (Hanwoo vs American vs Australian) had fatty acid profiles related to their diet during finishing (grain vs grass). They discovered noticeably higher unsaturated fatty acids in beef fed grain-based diets, especially in monounsaturated fatty acid (MUFA) content. In the United States, dietary inclusion of grain by-products has been implemented as a result of ethanol production utilizing corn grains for fuel production. As a result, energy dense distillers grains are often used as a substitute for part of the corn being used in livestock production.

Distillers grains are produced in a variety of forms, such as dried, wet, modified, or with soluble supplementation. All forms tend to be more energy dense than corn for livestock feed (Klopfenstein et al., 2008). As with corn, cattle fed diets with distillers grains inclusion deposit larger amounts of unsaturated fatty acids compared to grass-fed diets. In addition, distillers grains cause greater unsaturated fatty acids in meat compared to diets composed predominantly of corn (Nade et al., 2012; Chao et al., 2017; Ribeiro et

al., 2018). Fatty acid composition is a critical factor to meat quality, as unsaturated fatty acids are susceptible to oxidative mechanisms which impact lipid oxidation, color stability of lean proteins, and oxidative rancidity and off flavors, reducing consumer acceptability (Wood et al., 2004). de Mello et al. (2018) demonstrated greater inclusion of wet distillers grains compared to a corn-based diets increased malonaldehyde (MDA) content in whole muscle cuts, with MDA content increasing with increased days of retail display.

In sustaining meat quality, inclusion of vitamin E (α -tocopherol) is commonly utilized as a ROS scavenger deposited within organelle membranes. While α -tocopherol inclusion is more costly, it is suggested to improve immune function and weight gain in cattle production (Wood et al., 2004; Deters and Hansen, 2019). Also, inclusion of α -tocopherol has been shown to enhance meat quality (Faustman et al., 1998). Asghar et al. (1991), Ponnampalam et al. (2012), and E. N. Ponnampalam et al. (2012) showed greater stability of desirable meat color and decreased lipid oxidation in whole muscle products when animals diets were supplemented with α -tocopherol, in particular with greater inclusion of grass-based diets compared concentrates. Sales and Koukolová (2011) demonstrated how varying levels of α -tocopherol imparted within muscle related to color stability. Interestingly, the level of α -tocopherol deposited into meat eventually plateaus as vitamin E inclusion increases. These mirror redness (a^*) values in meat across fresh beef and pork products. This suggests that while the utilization of vitamin E in diets can greatly increase color sustainability, a limit of maximum efficacy is possible, so over supplementation of this feed additive may not be cost effective.

In addition, Chao et al. (2017) examined how cattle supplemented vitamin E in high energy diets can counteract oxidative stability in sarcoplasmic reticulum, reducing oxidative potential against organelle membranes and apoptotic mechanisms *in vivo* and early postmortem. This evidence confirms that the feeding strategies provided to livestock can have a direct impact of oxidative potential to affect meat quality.

Heat stress

Environmental factors have been suggested to increase oxidative factors. In particular, heat stress is a non-specific physiological response to prolonged exposure to high ambient temperatures (Xing et al., 2019). As a whole, heat stress is shown to reduce feed intake, as metabolic heat of digestion coupled with environmental heat promotes added stress on livestock (Habibian et al., 2016). It was suggested by Akbarian et al. (2016) that increased heat stress could create an imbalance of body redox stability, resulting in susceptibility to oxidative stress. Slimen et al. (2014) noted that heat stress caused an acute oxidative stress response, as heat stress upregulates superoxide anions and thermal inactivation of superoxide dismutase. Concurrently, heat stress promotes dysregulation of uncoupling proteins along the electron transport chain, allowing protein leakage and mitochondrial dysfunction (Akbarian et al., 2016). Interestingly, heat stress has been linked to variable response of oxidative stress, based on muscle type. Montilla et al. (2014) isolated red and white muscle fibers of pork semitendinosus muscles from gilts under 1 or 3 days of thermoneutral or heat stress conditions. As a whole, red muscle fibers expressed greater malonaldehyde (oxidation) across 1 day of heat stress, along with increased response to ROS enzymes and anti-apoptotic factors compared to

thermoneutral and heat stressed white fibers. This infers that red (oxidative) fibers are more sensitive to heat-mediated oxidative stress response systems and are most sensitive to downstream effects of oxidative stress. It can be postulated that prolonged-acute oxidative stress can be more detrimental to oxidative muscle fibers (Type I) compared to glycolytic white fibers (Type II).

Lipopolysaccharides

Given the complexity of oxidative stress *in vivo*, administration of pro-inflammatory stimuli are controlled measures of oxidative stress to monitor immune response factors. Lipopolysaccharides (LPS) are covalently-bound lipids and polysaccharides derived from the outer membrane of gram-negative bacteria (i.e. *Escherichia coli*) and are a known endotoxin used in acute inflammatory-mediated oxidative stress studies (Sternberg, 2007; Powers et al., 2011). As an endotoxin, LPS stimulates toll-like receptor signaling pathways, which initiate innate immune response mechanism NF κ B, to start transcription of pro-inflammatory cytokines interleukin-1 β (IL-1), interleukin-6 (IL-6), and tumor necrosis factor- α (TNF- α) (Halawa et al., 2013). Under condition of high LPS concentration, production of pro-inflammatory mediators such as ROS can be produced as a result of LPS toxicity, promoting oxidative stress to occur. Implementation of management strategies has been shown to influence the degree of LPS-mediated oxidative stress on livestock performance. Bai et al. (2019) illustrated a decrease in rate of gain in Cherry Valley meat ducks as LPS dosage increased.

Concurrently, an increase in inflammatory cytokines IL-6, IL-10, TGF- β , as well as an increase in oxidative synthase protein inducible nitric oxide synthase (iNOS) have

been found. This makes sense, as an increase in compensatory weight of liver, a metabolic organ, also occurred. Yates et al. (2011) reported LPS to increase inflammatory response hormone cortisol, rectal temperatures, and lymphocytes as mature ewes consume higher crude protein diets. Studies have been performed to investigate the effects of dietary oils on aggregated inflammatory conditions. Yang et al. (2008) compared the inclusion of fish or corn oil at 4.5% (g/kg of BW) on performance of chickens when administered an inner-perineal LPS challenge or saline control. Lipopolysaccharide challenge lowered overall feed efficiency and increased lymphocyte generation. Interestingly, inclusion of fish oil increased activity of inflammatory cytokines IL-1 compared to corn oil, and upregulated IL-6, and TNF- α compared to corn oil when exposed to an LPS challenge. In addition to pro-oxidative factors in diet, inclusion of exogenous antioxidants can ameliorate LPS-mediated oxidative stress. Kaiser et al. (2012) evaluated mRNA expression of different inflammatory cytokines in response to vitamin E type (natural vs synthetic) compared to level of intramuscular LPS injection. As expected, LPS injections resulted in higher RNA expression of cytokines compared to saline groups. Intriguingly, vitamin E source influenced different RNA expression of cytokines, as synthetic vitamin E increased IL-6 expression across saline groups and natural vitamin E increased TGF- β expression. Concurrently, natural vitamin E lowered iNOS expression compared to synthetic vitamin E, suggesting supplementing natural vitamin E inclusion may ameliorate inflammatory mechanisms *in vivo*.

Lipopolysaccharide-mediated oxidative stress, it is postulated to have an impact on muscle cells, which could affect meat quality postmortem. Suliman et al. (2004) examined gene expression of mitophagy and biogenesis in response to LPS-induced mice

harvested 6, 24, or 48 hours post-injection. Total oxidation (malondialdehyde; carbonyl content) increased as post-injection time of LPS increased coupled with significant decrease in reduced glutathione (antioxidant) content at 6 and 24 hours. Antioxidant activity aligns with expression of COX-1, an inflammatory producing enzyme, as COX-1 was at its highest expression at 0 and 6 hours, with a significant decrease after 24 hours post-injection. As post-injection time increased, there was a significant decrease in mtDNA expression 24 hours post-injection, which was negatively correlated with mitochondrial glutathione content ($r^2=0.89$) and positively correlated with malondialdehyde content ($r^2=0.91$). Concurrently, mitochondrial transcription factors Nrf-1 and Nrf-2 increased expression between 6 and 24 hours, but both ultimately decreased at 48 hours postmortem. This infers the lack of transcription factors for mitochondrial biogenesis occurred as post-injection time increased. Additionally, transcription factors used to generate protein complexes of the electron transport chain (ND-1 and ND-2) significantly decreased at 24- and 48-hours postmortem. Finally, an increase in uncoupling proteins was expressed as post-injection time increased, signifying the dissipation of the inner mitochondrial membrane. In total, the use of LPS caused an early antioxidant response, but quickly exhausted antioxidant effects. Coupled with the gradual decrease in proteins related to mitochondrial biogenesis, the induction of LPS-mediated oxidative stress could act as a stimulant for mitophagy. As a result, the onset of LPS-mediated oxidative stress could impact postmortem quality of meat by induced apoptotic mechanisms and oxidation of muscle proteins and lipids.

Tenderness

Since the phenomena of oxidative stress was discovered, it has been extensively linked to its effects on chronic inflammation and illness, and has been investigated thoroughly in medical literature. However, oxidative stress is now being linked to factors related to meat quality, as organelles linked to oxidative stress have been shown to impact proteolytic mechanisms related to postmortem quality of meat, influencing both product quality and consumer palatability. An investigation into oxidative stress on these organelles and mechanisms have been reported to hold an immediate impact on meat tenderization early postmortem (Cook et al., 1998; Wang et al., 1998; Koh and Tidball, 2000; Warner et al., 2005; Blunt et al., 2007; Cottrell et al., 2008; Archile-Contreras and Purslow, 2011; Senaratne, 2012; Mohrhauser et al., 2013; Ouali et al., 2013; Picard et al., 2014; Chao, 2015; Cottrell et al., 2015; Chao et al., 2017; Kunze et al., 2017; Chao et al., 2018; Liu et al., 2018; Wang et al., 2018; Zhang et al., 2018; Malheiros et al., 2019). Ouali et al. (2013) commented on how early postmortem tenderization in meat tends to show increased fractionation of cytochrome C as a result of greater meat tenderization. Cytochrome C fractions, along with other apoptotic precursors involved with the TCA cycle (3-Hydroxyisobutyrate dehydrogenase) increase early postmortem. In addition, Picard et al. (2014) reported how proteolytic enzymes (μ -calpains) and proteins related to muscle structure (myosin chains) are all highly active in meat during aging. These studies imply that apoptotic mechanisms can act as initiators of proteolytic enzymes linked to meat tenderization. It has been recognized, however, that meat given similar aging conditions with drastically different tenderness values exhibit unique proteomic profiles, which can insinuate how inherent differences in biological mechanisms can influence

meat tenderization. Malheiros et al. (2019) investigated differences in oxidized proteins from angus-crossbred cattle under identical management conditions, aged for two days postmortem, with different shear force values to designate between groups labeled tender ($38.2 \pm 2.9\text{N}$), intermediate ($51.9 \pm 6.8\text{N}$) and tough ($74.5 \pm 7.8\text{N}$). Amongst these groups, it was evident that tender samples had greater oxidative damage to structural (actin, myosin, desmin), regulatory (troponin), and antioxidant proteins (peroxiredoxin), along with some anti-apoptotic heat shock proteins compared to tough samples. Interestingly, tough samples exhibited less oxidation towards metabolic proteins, including enzyme SOD2 related to mitochondrial ROS stress. These data depict how early postmortem tenderization can be influenced by the expression or activity of oxidative stress factors, and how these factors can impact the oxidation of proteins which promote tenderization. Further investigation is necessary to discern how different ROS/RNS impact components of meat tenderness.

Both *in vivo* and *in vitro* analysis of muscle tissue has been examined. Using nitric oxide as the contributor to oxidative stress, Cook et al. (1998) induced pre-rigor *longissimus lumborum* with promoters and inhibitors of nitric oxide to assess postmortem meat quality. Across multiple days aging, Cook et al. (1998) discovered increased tenderness on 3 and 6 day aging in samples administered with NO promoters compared to NO inhibitor group. There were no tenderness differences in 1 or 8 days of aging. This data supports that *in vitro* NO generation in muscle can stimulate increased postmortem tenderization. However, this data is shown to be contrary compared to more recent literature. Koh and Tidball (2000) induced skeletal muscle cells with nitric oxide donors to inhibit the activity of m-calpain on the breakdown of cytoskeletal protein talin. These

data suggest that nitric oxide inhibits m-calpain activity by S-nitrosylation of the cysteine sites on the endogenous enzyme. In addition to calpain inhibition, the data shows upregulated calpastatin activity, further inhibiting calpain-mediated proteolysis of structural proteins. Liu et al. (2018) and Warner et al. (2005) theorize reduced tenderization is a result of nitrosylation reactions of NO with organelle calcium channels RyR1 and SERCA. Due to NO nitrosylation, there is insufficient cytosolic calcium available to facilitate calpain autolysis on cytoskeletal proteins such as titin, desmin, or nebulin, resulting in less tender meat. Conversely, inhibition of nitric oxide has been indicated to increase postmortem tenderization. Cottrell et al. (2008) indicated antemortem injections of nitric oxide inhibitors 135 minutes prior to slaughter can improve meat tenderness early postmortem in lambs. Combining NO inhibitors with brief exercise prior to slaughter, lambs had a lesser degree of pH decline during rigor mortis, although gross numbers are not significant enough to be a detriment to meat quality. Interestingly, Cottrell et al. (2008) evaluated across two muscles: *longissimus lumborum* (LL) and *semimembranosus* (SM). Interestingly, NO inhibition was shown to significantly decrease shear force in LL muscle ($p < 0.01$), with no change in shear force of SM ($p = .48$). This data was corroborated when Cottrell et al. (2015) exposed sedentary lambs to NO donors and inhibitors 190 minutes prior to slaughter. Cottrell et al. (2015) exhibited that SM tended to increase in shear force at 1-day aging when administered NO inhibitors compared to NO promoters ($p = .06$). While not significant ($p = .58$), NO inhibitors lowered shear force in LL muscle across 1 and 3 days of aging compared to NO promoters and control. While NO inhibitors did change the progression of pH decline

in SM muscle, the ultimate pH was equivalent to other treatments and did not alter meat color characteristics across treatment or muscle type.

From these different studies it can be suggested that NO interaction may be more muscle dependent than originally considered. In conjunction with shear force, proteolytic activity can change as a result of RNS. Zhang et al. (2018) investigated changes in protein proteolysis at 1-day postmortem across samples incubated with varying degrees of either NO donor or inhibitor. Nitric oxide inhibitor treatments exhibited less total protein nitrosylation and greater autolysis of calpains, desmin, troponin-T, myosin, and protein solubility compared to NO donors treatments and control groups, indicating greater proteolysis. There was a significant interaction between the treatment and aging time, with the increasing amount of NO inhibitor resulted in greater protein degradation over time compared to all other NO donor treatments. Through this data, it is conceivable to state NO may play a critical role in meat quality through regulating calpain autolysis and myofibrillar protein degradation during postmortem aging.

In addition to NO, isolation of H₂O₂ has been evaluated to influence meat tenderness. It is speculated that the function of H₂O₂ may be due to its amount within the muscle cell. Archile-Contreras and Purslow (2011) demonstrated how H₂O₂-mediated oxidative stress can influence collagen turnover in different muscles. Given collagen is a structural protein of various connective tissues throughout muscle, its presence can impact on overall palatability of meat, including tenderness. Incubating myofibrillar muscle cells from 1-hour postmortem *longissimus dorsi* and *semitendinosus* muscle, they introduced samples to either 0.5 or 5 μ M of either H₂O₂ or synthetic pro-oxidant xanthine oxidase. In particular, an inverse relationship between H₂O₂ concentration and matrix

metallopeptidase-2 (MMP-2) growth, with a significant increase in MMP-2 growth in *longissimus dorsi* muscle compared to *semitendinosus*. MMP-2 is a matrix metalloproteinase used to facilitate collagen turnover. Additionally, increased H₂O₂ concentration reduced total soluble collagen synthesis, with a noticeable loss of collagen synthesis in *semitendinosus* muscle. As a result, they were able to relate different levels of ROS production to the degree of collagen turnover across different muscles, with the overall indication that ROS reduced collagen synthesis. As collagen is a contributor to beef tenderness, this study reinforces the impact of ROS factors and their implications towards meat quality and muscle structure postmortem. As Blunt et al. (2007) discovered, hydrogen peroxide incubation with ischemia rat ventricular muscle is shown to reduce desmin degradation. Instead, H₂O₂ is theorized to activate MAPK pathways for cell sustainability. This is validated by showing decreased desmin degradation in conjunction with incubating myocytes with H₂O₂ and a MAPK inhibitor. Furthermore, Blunt et al. (2007) depicted increased expression of HSP27 on decreased desmin degradation regardless of calpain activity. This suggests that H₂O₂ may be a driving factor of HSP27 expression early postmortem, preventing calpain activity and reducing overall protein degradation. Interestingly, the analysis presented H₂O₂ as a promoter of protein degradation during increased calcium availability, as H₂O₂ combined with calcium had numerically lowered desmin degradation. These amounts of degradation were statistically significant when H₂O₂ and calcium were combined with MAPK inhibitors. This is telling, as these conditions best represent cellular stability during apoptosis via MAPK inactivity, oxidative stress factor H₂O₂, combined with increased cytosolic calcium as an indicator of cellular death postmortem. Given the combination of these conditions, it is significant

when considering all these factors work in concert of one another during cellular death cascades as a result of oxidative stress, and a significant source of calcium early postmortem could provide the tools necessary to facilitate increased protein degradation. Wang et al. (2018) investigated this idea: the effect of reactive oxygen species-mediated oxidative stress on mitochondrial apoptosis factors in relation to tenderness in yak meat. By injecting 20mM of either H₂O₂ or H₂O₂-inhibitors compared to a control group across multiple aging periods, it was discovered that H₂O₂ increases total ROS, apoptosis regulator proteins (Bax), lipid oxidation, and caspase activity compared to both the control and inhibitor treatment. Concurrently, H₂O₂ decreases SOD, glutathione peroxidase, mitochondrial stability via the mitochondrial permeable transition pore opening, anti-apoptotic proteins (Bcl-2), and shear force in yak meat throughout a 7-day aging period. This data supports the theory of oxidative stress and its influence on mitochondrial stability, calcium release, initiating apoptotic mechanisms. These observations further support the idea that oxidative stress has a role in the activation of mitochondrial apoptosis and meat tenderness by influencing regulators necessary of the apoptotic pathway. Furthermore, Wang et al. (1998) and Mohrhauser et al. (2013) decipher that caspase, while an indicator of mitochondrial apoptosis, is not the primary cysteine enzyme used to inhibit calpastatin activity. Rather, μ -calpain is the primary inhibitor of calpastatin in conjunction with caspase enzymes. It also reinforces that calpains are the primary protease responsible for myofibrillar protein degradation during postmortem aging of myofibrils.

As suggested, diet has been used as a possible indicator of changes in tenderness in beef cattle research. Recent work at the University of Nebraska-Lincoln has identified

an improvement in tenderness when distillers grains are supplemented (Senaratne, 2012; Chao, 2015; Chao et al., 2017; Kunze et al., 2017; Chao et al., 2018). Senarte (2012) indicated that meat from cattle fed distillers grains, without supplement of vitamin E, increased both tenderness and troponin-T degradation compared to steaks from cattle fed a corn-based control diet. These studies suggest that muscle membrane integrity can be altered and compromised as a result of feeding distillers grains, which could promote greater calcium release early postmortem and increase calpain activity to promote improved tenderness. Knowing this, Chao (2015) indicated an increase in PUFAs in the sarcoplasmic reticulum which could lead to greater calcium leakage early postmortem, upregulating increased enzymatic tenderization. Kunze et al. (2017) investigated further into this occurrence and isolated mitochondria to understand the calcium flux mechanism in relation to beef steak tenderness. This study illustrated that cattle fed distillers grains deposited greater amounts of linoleic acid (18:2) in addition to total PUFA content in mitochondria. Concurrently, cattle fed predominantly corn diets had mitochondria which tended to retain more calcium compared to cattle finished on distillers grains ($p = 0.08$). Therefore, it can be speculated that greater PUFA content in organelles can increase susceptibility of oxidation reactions, altering calcium flux and resulting in increased tenderness early postmortem. Chao et al. (2018) indicated that distillers grains may impact sarcoplasmic reticulum composition as well, as feeding distillers grains increased total PUFA in sarcoplasmic reticulum membrane. In turn, increased unsaturated fatty acid content could lead to greater oxidative potential, facilitating calcium leakage to improve tenderness early postmortem. These studies strengthened the concept of which oxidative stress factors can alter biological composition, and prolong oxidative stress prior to death,

while not becoming detrimental to animal well-being. This could be a major facet to enhance meat tenderization and produce more consistently tender meat products moving forward.

Color

Color is the primary purchasing factor consumers use when evaluating meat quality (Suman et al., 2014). The demand to maintain ideal fresh meat color is an ongoing challenge, as industry implemented techniques to enhance product palatability can greatly influence the stability of meat color during retail display. Of these techniques, tenderization of meat during aging reduces prolonged color stability in meat, as the substrates used in redox reactions are slowly depleted as aging progresses. Mitacek et al. (2019) reported a decrease in color stability during increased aging periods (3 to 28 days aging). Factors such as oxygen availability, mitochondrial activity, and color redox-reaction substrates all contribute to color stability. Oxygen depletion increased as aging continued, in addition to lower mitochondrial concentrations. Metabolomic analysis determined a decrease in substrates necessary to run TCA cycle, supporting lack of mitochondria activity. Interestingly, Mitacek et al. (2019) discovered greater NADH reductase activity with increased aging, meaning an increase in demand of NADH redox reactions are necessary to sustain oxymyoglobin pigment as meat ages. Concurrently, total NADH concentration decreased as aging increased, and an increase in oxidative potential as indicated by increased malonaldehyde as aging and days of retail display interaction. These results indicate that there are insufficient substrates necessary to supply the growing requirements of redox reactions to maintain color stability in aged meat,

allowing further oxidation of meat tissue using pro-oxidants as means for oxidation, including factors related to oxidative stress. In addition to aging effects, muscle type has been indicated as a contributing factor to color stability. Ke et al. (2017) discovered greater rate of decline in a^* (redness) values and increased metmyoglobin content in *psoas major* (PM) muscle compared to *longissimus lumborum* (LL) of select graded beef cattle. Psoas major muscle had a greater rate of decline of oxygen consumption, mitochondrial activity, metmyoglobin reducing activity, and lipid oxidation compared to LL. Interestingly, PM expressed greater cytochrome C across all retail display days compared to LL, indicating an increased rate of cellular degradation. This data can speculate that an increase in mitochondrial degradation caused by oxidation-mediated apoptosis, as evidenced by cytochrome C expression, can influence meat color stability. This relationship can be a contributing factor when meat is introduced to oxidative stress or its precursors.

When examining oxidative stress inclusion antemortem, however, there is minimal effect on meat color. Ponnampalam et al. (2005) discovered no differences between color stability of *longissimus thoracis lumborum* (LTL) or *semimembranosus* (SM) when administered nitric oxide donors and inhibitors into lambs twenty-four hours prior to slaughter. Concurrently, Cottrell et al. (2008) found no differences in color measurements on LTL or SM across 1- and 3-days aging when animals were induced with a NO inhibitor prior to slaughter. A follow up study by Cottrell et al. (2015) indicated that lambs administered NO donors 190 minutes prior to slaughter exhibited no significant changes in color of LTL or SM muscles across 1 and 3 days of aging. Concurrently, Niu et al. (2016) used a 3-day LPS challenge across two LPS

concentrations (3 mg/kg; 6 mg/kg) to evaluate meat quality of broilers breast and thigh muscle against a saline control. Both muscles had a lower ultimate pH 24 hours postmortem as LPS increased, however, the gross differences between control, 3, and 6 treatments were negligible. In both breast and thigh muscle, an increase ($p < 0.05$) in b^* values on the 3rd day of retail display was found as LPS concentration increased, with breast muscle L^* values increasing ($p < 0.05$) as LPS increased on the 6th day of retail display. While these color values were significant, however, their gross differences across treatments were negligible. It is speculated that a more prolonged oxidative stress challenge may be more critical to understand its relationship to meat color.

Protein oxidation

As protein oxidation occurs, the modification of muscle tissue from carbonylation is shown to affect muscle and color (Batifoulier et al., 2002; Santé-Lhoutellier et al., 2008; Terevinto et al., 2010). Promeyrat et al. (2011) used this proteomics to build correlations of carbonyl formation and protein oxidation in m. *Longissimus lumborum* of pigs across different storage periods. Notably, they showed a negative correlation between carbonyl content and activity of superoxide dismutase activity. This indicates that by reducing the formation of free radicals, SOD acts as regulators of protein stability against oxidation. There also was a negative correlation between myoglobin and carbonyls at 4 days of storage. This contrasts with 1-day storage positively correlating the two variables. This contrasting data infers myoglobin to act as both an antioxidant and pro-oxidant, dependent on its current state. In the presence of hydrogen peroxide, metmyoglobin is transformed into perferryl-myoglobin, which can be autoreduced by

oxidative radicals at low pH (~5.5) which falls in line with the ultimate pH (pH_u) of fresh meat (~5.4-5.8). Given the capacity for perferryl-myoglobin to reduce oxidative radicals, in conjunction with prolonged postmortem storage, metmyoglobin formation could promote activity of oxidant enzymes such as catalase and glutathione peroxidase.

It has been postulated increased carbonyl formation may be an indicator for meat texture, as Carlin et al. (2006) discerned proteolytic activity of calpains to be hindered by carbonylation, decreasing postmortem tenderness. Due to the rich concentration of cysteine residues comprised within the calpain enzymes, oxidative inactivation of such amino acids could hinder proteolytic function. If protein oxidation mechanisms occur during aging, cross-linking of proteins could help explain the correlation between carbonyl content and decreased tenderness in meat (Lund et al., 2007). Carbonylation has been indicated as a result of different feed strategies. Santé-Lhoutellier et al. (2008) demonstrated increased carbonyl formation in concentrate (corn-based) diets compared to pasture-fed in lamb during prolonged meat storage. Carbonyl content could not explain protein digestibility of myofibrillar proteins. Conversely, Batifoulier et al. (2002) demonstrated diet inclusion of vitamin E to retard the loss of free thiols in microsomal organelles in turkey breast muscle. In addition, turkeys fed vitamin E exhibited less malonaldehyde content across days of storage. Building off this data, Terevinto et al. (2010) evaluated the oxidative stability and endogenous enzyme response in three muscles of rhea, a species closely related to ostrich. They discovered muscle exhibiting higher malonaldehyde content also had greater carbonyl content across muscles, demonstrating that there may be inherent differences between muscles and their capacity to generate carbonyls. Given the differences in color stability of different muscle

combined with the interrelationship between color and lipid oxidation, it is plausible that different muscle are more susceptible to oxidation (Faustman et al., 2010).

Lipid oxidation

Given the link between oxidation of fatty acids and oxidative stress, evaluation of lipid oxidation is critical when examining meat quality. As shown by Gatellier et al., (2004), Zhang et al. (2011), Chao et al. (2018), and Zhao et al. (2018), diet can impact lipid and total oxidation, as an increase in unsaturated fatty acids and oxidized oils can increase the oxidative potential postmortem. Interestingly, Zhao et al. (2018) evaluated the generation of ROS in conjunction with lipid oxidation when lambs were supplemented grape pomace. They noted a decrease ($P < 0.05$) in total ROS as inclusion of antioxidants increased. Concurrently, there was a significant decrease ($P < 0.05$) in malonaldehyde content as antioxidant content increased. In addition, evaluating the onset of oxidative stress was used without dietary treatments. Niu et al. (2016) evaluated LPS-mediated oxidative stress on oxidative stability of broilers. Malonaldehyde content did increase ($p < 0.05$), as well as a decrease in total antioxidant capacity, in plasma as LPS concentration increased. Furthermore, malonaldehyde content increased ($p < 0.001$) as LPS increased across all days of retail display. Additionally, Pradhan et al. (2000) used endogenous catalase as a measure of lipid oxidation in muscle tissue. They discovered that increased catalase inclusion in ground beef semitendinosus decreased lipid oxidation across all storage periods, signifying a decrease in oxidative potential postmortem due to the increase in endogenous enzymes.

Evaluation of oxidative stress markers

In order to better understand oxidative stress and its link to muscle cell interactions and meat quality, there is an ever-growing demand to understand its behavior from the onset of ROS generation, its relationship with physiological response mechanisms, and the activity of prolonged oxidative stress on muscle characteristics both *in vivo* and postmortem. However, the evaluation of oxidative stress factors, and their intermediate constituents, is a challenge. Due to the complexity of oxidative stress and the short half-life of many constituents, strategies used to validate its effects on meat quality have their constraints. In the literature, there are a multitude of oxidative stress products produced in muscle tissue which can be used as markers indicating oxidative stress has occurred, and relative quantities of such compounds are being linked to meat quality attributes (i.e., carbonyls) (Lawson et al., 1999; Dalle-Donne et al., 2003; Ponnampalam et al., 2017). However, the obvious constraint of this approach is the lack of knowledge on the degree of oxidative stress occurring *in vivo* and the regulation of different pathways oxidative stress activates on a per organism basis, as all can have variable levels of physiological response mechanisms. In the last two decades, there is growing interest for *in vivo* evaluation of oxidative stress factors to monitor their quantification and sustained activity during stages of oxidative stress. This method of analysis is even more complicated, due to the inherent behavior of ROS and their average half-lives (~2-5 seconds) (Bekhit et al., 2013). Additionally, there is still developing technology sensitive to evaluate real-time analysis of ROS activity. In order to better understand the totality of oxidative stress, there is value to use both measures to create a more rounded picture of oxidative stress on muscle and meat quality.

In vivo: single wall nanotube hydrogels

Recent milestones in biotechnology have enabled the development of biosensors used to evaluate real-time generation of molecular ions. In particular, the use of single-walled carbon nanotubes (SWNTs) as an active biosensor has sparked interest in medical applications for its innovative methodology and growing accuracy to detect molecules in living organisms, such as glucose detection (Strano et al., 2004). In short, SWNT are a synthetically manufactured graphene cylinders made of a 1-D carbon atoms rolled into a cylindrical shape. Commercially, the most common SWNT used in biotechnology is the CoMoCAT (6,5) TM, as the binding of the carbon molecules acts as a very stable semiconductor. Their high conductivity facilitates stable emission of near-infrared (NIR) light to transmit through SWNT with no noticeable photo-intensity threshold, this facilitates prolonged imaging of SWNT in organic tissue and cells without degradation when introduced into organisms (J. Zhang et al., 2011). As a result, the sustainability of SWNT under intense electromagnetic radiation makes them a useful tool in molecular detection using real-time evaluation. During manufacturing, the composition of the SWNT is critical to its end function, especially considering both the length and diameter of the SWNT can influence the sensitivity and compatibility to bind with unique DNA sequences (J. Zhang et al., 2011). Iverson et al. (2015) illustrated wrapping SWNT with a 15-repetition of adenine-thymine complex [(AT)₁₅] as a sensor for nitric oxide.

Concurrently, the development of (GT)₁₅ DNA sequences have been indicated as reliable sensors for H₂O₂ detection (Kim et al., 2011). However, the (GT)₁₅ sequence is sensitive to both H₂O₂ and NO, requiring the use of both DNA-wrappings to detect H₂O₂ quantity via the transitive property (Iverson et al., 2017). In addition to SWNT

manufacture, the specificity and wrapping of DNA is theorized as a critical component to the sensitivity of detecting molecules. Unfortunately, techniques to DNA wrapping are still undefined, as this field of research is still in its early stages (Iverson et al., 2013). When administered in a biological system (i.e., the blood/tissue), there is speculation that SWNT can degrade from itself and independently traverse throughout cells. In order to contain SWNT within a localized system, mixed alginate gels have been used to contain SWNT upon administration into a biological system (Iverson et al., 2013). The binding of SWNT to the gels allows additional support for SWNT to stay bunched together, facilitating a better composition for SWNT upon reading fluorescence (Iverson et al., 2015; Iverson et al., 2017). When evaluating single-molecule components, excitation of DNA-wrapped SWNT occurs during exposure to a NIR stimulus laser, which elicits a fluorescent intensity of SWNT. This intensity can be used to dictate the total quenching of SWNT in the presence of the targeted molecules, with less fluorescence equating to greater quenching of SWNT, indicating a greater concentration of the targeted molecule present (Strano et al., 2004). Concurrently, consecutive readings of SWNT can be used to evaluate both the rate of quenching from the time of exposure to a stimuli, as well as the rate of recovery, the time in which the SWNT regains its fluorescence (Jin et al., 2010; Iverson et al., 2017). While still in its infancy, the field of carbon nanotubes is growing with exponential potential on scientific applications. If administered and measured properly, SWNT could be used as an innovative technique to evaluate the onset of oxidative stress and rate of physiological response for *in vivo* livestock analysis.

Ex vivo: omic profiles

When examining the development of muscle tissue, there are countless conditions which govern the growth, functionality, and stability of muscle cells, all of which contribute to the composition of meat quality after harvest. These variables are strongly influenced through both predisposed genetic development in addition to extrinsic conditions such as species, diet, pre- and post-slaughter handling, processing, and formulation of products (Capozzi et al., 2017). These factors trigger changes in how the organism, and its subsequent tissues, acts and reacts to certain stimuli. These factors can be used to distinguish differences in muscle cells through omics analysis.

A novel, comprehensive approach, omics are the analysis of genomic and molecular profiles of an organism (D'Alessandro and Zolla, 2013). Using this tool, examination of different components of muscle growth and function in relation to oxidative stress factors are possible. In meat science, transcriptomics has been used to relate molecular mechanisms underlying the formation of meat quality traits with gene expression (Guo and Dalrymple, 2017). In particular, the relationship of upregulating or downregulating specific pathways related to muscle cell proliferation, growth, organelle development, cell modification, enzymatic activation, and cell immunity are of interest. Bongiorni et al. (2016) and Arora et al. (2019) examined breed variation on the impact of muscle development and tenderization in cattle and sheep, respectively. These scientists used pre-existing data (tenderness, cell proliferation) as benchmarks to assess genomic differences between treatments (breed). Guo and Dalrymple (2017) utilized transcriptomics to analyze biological mechanisms which regulate lipogenesis, fatty acid deposition, and tenderness characteristics related to intramuscular fat in livestock species.

Conversely, transcriptomics can be used to assess genomic conditions in response to specific stimuli, such as oxidative stress. Kim et al. (2018) used oxidative stress as a treatment to assess the expression of genes in specific tissues of mice. As expected, an overall upregulation of ROS enzymes (catalase, superoxide dismutase, glutathione peroxidase) were found. Interestingly, the degree of upregulated genes was tissue-dependent, as metabolic organs (liver and kidney) were greater in total expression compared to muscle tissue. This investigation revealed gene expression can be dependent on the tissue of interest, and that each tissue may respond to a treatment with varying magnitudes of gene expression. Transcriptomics acts as an outline, as the complex expression of different biological pathways elucidates how genetic mechanisms interact with one another to reach a final function.

Proteomics, the study of biological proteins, can assess the composition of muscle proteins under specialized conditions. This is appropriate, as muscle proteins can be used in association with different meat quality attributes, such as calpains for enzymatic tenderization. This association is indicative of the conditions which developed the tissue in its current state (i.e., genetics, environment). Hollung et al. (2007) identified individual proteins related to mechanisms responsible in postmortem tenderness, such as myofibrillar proteins and the calpain system, in addition to proteomes related to muscle color (sarcoplasmic) and antemortem stress (heat shock proteins and glycolytic enzymes). Building off of this research, Polati et al. (2012) distinguished proteomic changes as a result of proteolytic aging. As aging increased, the expression rate of troponin-T, α -actinin, and myosin light chains all increased, supporting the role of proteolytic breakdown of essential proteins to meat tenderness. Concurrently, a mixed expression of

heat shock proteins (HSP) and α -crystallin were shown to increase with prolonged aging. The expression of these proteins can be used as markers for meat tenderness, with aging as a key indicator in the change of protein expression and functionality postmortem. In addition, Picard et al. (2014) demonstrated genetic variation has an impact on meat quality, as differing cattle breeds show varying expression of metabolic (lactate dehydrogenase) and contractile (myosin, actin) proteins in relation to different muscles within a breed. This suggests that composition of muscle fibers, in addition to metabolic properties of muscle cell, can be dependent on genetic predisposition (Picard et al., 2014; De Souza Rodrigues et al., 2017). When referring to meat tenderness, the common comparison between *Bos indicus* and *Bos taurus* cattle is utilized, as both species have distinctly different meat quality attributes. Proteomic evaluation reveals *Bos indicus* cattle presented higher expression of myosin and actin proteins, enhancing contraction force in skeletal muscle, in addition to increased expression of heat shock proteins 5 (HSP5) located within the sarcoplasmic reticulum. Concomitantly, *Bos indicus* cattle held a greater proportion of fast-twitch glycolytic fibers, compared to the oxidative slow-twitch fibers of *Bos taurus* cattle. Conversely, *Bos taurus* exhibited higher phosphorylation in troponin-T and HSP9, both of which have been linked to increased μ -calpain activity (De Souza Rodrigues et al., 2017). Couple these proteins with greater shear force (less tender) values in *Bos indicus* cattle, and it can be illustrated how a genetic bias can influence inherent quality traits in fresh meat products.

Metabolomics is used for the quantification of cellular molecules taking part as intermediates or end products of metabolic reactions (Capozzi et al., 2017). The identification of metabolic substrates can be used as a reference in discerning subtle

differences in a given set of conditions. Straadt et al. (2014) used this method to evaluate sensory attributes in thawed pork chops of crossbred pigs related to the composition of amino acids extracted during drip loss. While the flavor of meat is commonly associated with cooked meat via the maillard reaction, the composition of non-volatile compounds in raw meat such as amino acids, can elicit the unique flavor profile (Mottram, 1998). These amino acids, while readily available in post-rigor meat, have been shown to increase in concentration during aging, and impact a greater depth of flavor as a result (Nishimura et al., 1988). Additionally, Kim et al. (2016) used dry aging as a model to enhance amino acid concentration of glutamate, tryptophan, and isoleucine, which are associated with the umami, bitter, and maillard flavor profiles, respectively. The development of dry aging has been suggested to elicit greater complexity of flavor being present compared to wet aged beef, which builds its niche market for this unique flavor profile. However, further investigation into amino acid composition is needed to interpret its impact of meat flavor. Chen and Ho (2002) have identified offsetting flavor attributes of amino acids. Carnosine, which has been linked to “roasted” and “nutty” notes in cooked meat, has also been implicated to reduce compound 2-methyl-3-furanthiol, a cysteine based chemical responsible for the “meaty” aroma consumer identify in cooked meat products. A greater demand for metabolomic analysis is necessary to tackle the current challenges to relate specific amino acids to flavor profiles of aged meat, increasing the viable application of this methodology.

Isoprostanes

In addition to omics measurements, the evaluation of other downstream end products *specific* to oxidative stress have been suggested. While oxidative stress factors can induce oxidation of both lipids (malonaldehyde) and proteins (carbonyls/thiols), the isolation of F₂-isoprostanes has been recognized as a key indicator for oxidative stress (Basu, 1998; Lawson et al., 1999; Montuschi et al., 2004; Nikolaidis et al., 2011). Compared to other oxidation products, generation of F₂-isoprostanes cannot be accomplished via photo-oxidation or auto-oxidation (Lawson et al., 1999). Rather, an integrated mechanism is required to produce isoprostanes, which is solely contingent on ROS-mediated oxidation (Lawson et al., 1999; Milne et al., 2011). Currently, F₂-isoprostanes are the family of prostaglandin-like compounds formed by non-enzymatic, free-radical oxidation of arachidonic acid (20:4) with reactive oxygen species. During oxidation, the abstraction of hydrogens due to ROS result in delocalized pentadienyl carbon-centered radicals (Lawson et al., 1999; Montuschi et al., 2004; Nikolaidis et al., 2011). Subsequently, peroxy radicals formed by oxygen molecule binding undergo cyclization, the formation of carbon rings, using additional oxygen molecules. These molecules then attach hydroxyl radical groups (·OH) as a final binding reaction, resulting in isoprostanes formation. Within the F₂-isoprostane group, different isoforms are produced based on which carbon the allylic hydroxyl group binds (Figure 6) (Nikolaidis et al., 2011). Generated during lipid peroxidation, F₂-isoprostanes are produced as esterified fatty acid form in phospholipids membranes, and then released using phospholipase action (Montuschi et al., 2004). Stafforini et al. (2006) and Milne et al. (2011) establish phospholipase action as digestion enzymes, facilitating the breakdown of

phospholipid arachidonic acid. Upon release, F₂-isoprostanes can follow oxidized fatty acids into the blood, or transfer out from the kidneys and into urine. Compared to other oxidative products, F₂-isoprostanes very stable compounds that are detectable in all normal biological fluids and tissues. When evaluating F₂-isoprostanes, different collection methods can discern variation in detected levels of F₂-isoprostanes. Urine sampling can detect unesterified (free) fatty acids which can transport into the urine, which is easily collected. Both blood plasma and skeletal muscle still contain esterified F₂-isoprostanes, which suggest the quantity of F₂-isoprostanes formed to be dependent on the quantity of lipids present, especially arachidonic acid (Nikolaidis et al., 2011). However, Basu (1998) noted blood plasma to quickly metabolize F₂-isoprostanes, with a half-life no longer than 20 minutes, suggesting blood plasma to have greater variability of F₂-isoprostanes when mishandled. However, immediate analysis of blood plasma F₂-isoprostanes provides an index of total endogenous production *in vivo*. Levels of esterified F₂-isoprostanes in muscle can be used to localize oxidation in an area of interest (skeletal muscle). This is common to evaluate for muscle biopsies or sampling in anoxic muscle tissue, as the lack of blood prevent F₂-isoprostanes to be transferred and metabolized in blood.

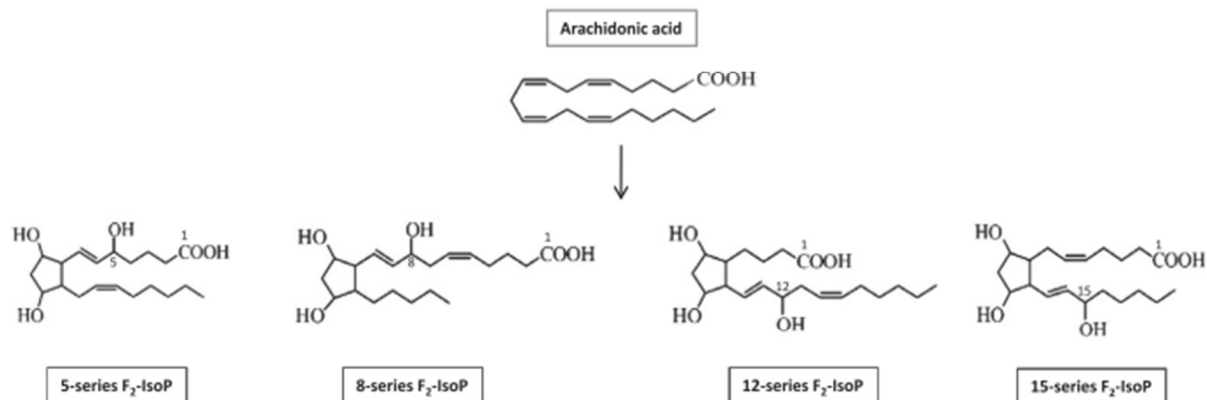


Figure 6: Chemical structure of arachidonic acid and the four regional isomers of isoprostanes (Nikolaidis et al., 2011).

After collecting a sample, there are a multitude of methods used to detect isoprostane quantity. Lawson et al. (1999) Roberts and Morrow (2000) illustrate the use of mass spectrometry (MS), as this was the original and ideal methodology used to evaluate isoprostanes. Combine this with liquid/gas chromatography (LC/GC), and F₂-isoprostanes detection is highly specific and sensitive. However, this is a very expensive process which deters its use in contemporary evaluation of isoprostanes. Currently, the generation of enzyme-linked immunosorbent assays (ELISA) are frequently used as modes for isoprostane detection. Commercial ELISA kits can detect individual isoforms of F₂-isoprostanes. While ELISA kits are more cost-effective and hold respectable sensitivity to isoprostane detection compared to GC-MS/LC-MS (75-90% and >96%, respectively), the sensitivity of ELISA kits require high quality samples, which have gone through a purification method to eliminate any biological contaminants which could interfere with the immunoassays.

Given the relationship between isoprostane generation with oxidative stress activity, detection of this molecule is regarded as the standard for evaluating oxidative stress.

However, there is very little literature relating this oxidative stress to meat quality using

this molecule as a baseline measurement. Karamouzis et al. (2004) used microdialysis probes to evaluate the generation of 8-isoprostane $\text{PGF}_2\alpha$, a predominant isoform of F_2 -isoprostanes, in the event of rest compared to prolonged exercise. They found that exercise increased ($P < 0.05$) isoprostane content nearly three-fold. This makes sense, as an increase in exercise promotes an upregulation of ROS generation (Li et al., 1999). Given the generation of oxidative stress on muscle tissue, Ponnampalam et al. (2017) theorized the change in isoprostanes as an *in vivo* indicator of meat quality deterioration postmortem (*in vitro*). Using diet as a treatment of metabolic energy, lambs were fed for 8 weeks on diets consisting of varying levels of metabolizable energy in feedlot, ryegrass, and lucerne hay (12.9, 10.4, 8.6 MJ/kg DM, respectively). Using blood samples, they detected an increase in plasma of 8-isoprostane $\text{PGF}_2\alpha$ levels, with the feedlot diet exhibiting the greatest isoprostane content at 4 ($P < 0.01$) and 8 ($P = 0.05$) weeks of feeding. This is due to both the increase in energy content of diets, but also the amount of α -tocopherol increased in roughage-based diets (ryegrass, lucerne). Positive correlations between 8-isoprostane $\text{PGF}_2\alpha$ levels and lipid oxidation (malonaldehyde content) have been found. Interestingly, positive correlations between 8-isoprostane $\text{PGF}_2\alpha$ levels and both arachidonic (20:4) and linolenic (18:2n-6) acid were significant ($P = 0.005$; $P < 0.001$, respectively). This supports Milne et al. (2011) on how F_2 -isoprostanes can potentially be generated from other PUFAs [eicosapentaenoic acid (20:5n-3), adrenic acid (22:4n-6), and docosahexaenoic acid (22:6n-3)] due to the orientation of their double bonds, but the knowledge on these fatty acids to generate F_2 -isoprostanes is not fully understood. As a result, the generation of F_2 -isoprostanes is an excellent marker for *in vivo* oxidative damage as a result of oxidative stress.

Other metabolic indicators

Physiological stress triggers a vast number of homeostatic mechanisms, producing signaling molecules and hormones to activate suppression of stress-inducing factors. In turn, a vast majority of these signaling factors are generated independent to a particular stress-induced event, allowing their evaluation as an overarching indicator of physiological stress. Cortisol, synthesized from cholesterol, acts to suppress stress by signaling other homeostatic pathways by binding to cortisol receptors, assisting in regulation of blood pressure, energy metabolism, and immune system response (Lee et al., 2015). Under conditions of oxidative damage, the upregulation of cortisol is an appropriate response. Hoogenboom et al. (2012) identified an influx in cortisol concentration in conjunction with testosterone release during spawning of brown trout (*Salmo trutta, L.*). This is appropriate, as reproduction is recognized as a benchmark for increased stress during an animal's life span. Under livestock management conditions, these benchmarks would include events such as birth, weaning, transportation, feeding strategies, and handling. All of these factors can be implicated as points of cortisol growth, and potential benchmarks for oxidative stress to occur.

In addition, the production of additional indicators is used to assess biological stress. Temperature is an ubiquitous measure of biological stress, as temperature increase is a strong correlation to acute illness or inflammation. Concurrently, increased environmental temperature can increase metabolic stress. Srikandakumar et al. (2003) indicated that increased temperature can stimulate an increase in rectal temperature, respiration rate, and alters blood chemistry in mature sheep. Interestingly, different breeds of sheep within the study has a lesser degree of overall stress (temperature, blood

chemistry) in response to increased temperatures, indicating that genetics may influence the capacity for animals to suppress biological stressors. As a whole, stress response studies can use baseline measurements as markers of biological stress ensuing in livestock.

Link between downstream effects of oxidative stress and meat quality

Components related to cellular respiration are critical to the development of ATP, organelle function for muscle contraction, and downstream effects towards postmortem tenderization of muscle (Kemp et al., 2010). Concurrently, the action of cellular respiration promotes the generation of ROS/RNS properties, and production of these compounds accumulate with physiological stress factors such as chronic inflammation or illness, even when these factors are not a recognizable detriment to the organism *in vivo*. If ROS/RNS generation overwhelms biological homeostatic response systems, oxidative stress begins, and is capable to manipulate a multitude of cellular proteins, lipid peroxidation, and regulation of nucleic information (Powers et al., 2011). As a result, organelles and their constituents related to muscle contraction and tenderization are distorted, with increased membrane leakage and overall organelle dysfunction. This cascade of events will alter both the composition and function of organelles, inciting cellular apoptotic pathways and development of new protein and lipid turnover. If oxidative stress occurs during death, especially due to chronic oxidative stress, the current state of cellular dysfunction may expose molecular ions readily available in the cytosol, as well as nucleic expression of postmortem mechanisms (Kemp and Parr, 2012).

Tenderness is one of the most important factors for consumer palatability and repurchasing of meat (Koochmaraie and Geesink, 2006; Kemp et al., 2010). However, subtle variations in tenderness within muscles have been shown to greatly influence consumer likeness to different retail products (Martinez et al., 2017). One method to mitigate inconsistent tenderness is producing high marbled cattle, as an increase in intramuscular fat is shown to increase overall tenderness and palatability scores in beef strip loins (Platter et al., 2003). However, these tenderness differences decrease with lower marbled meat, providing greater challenges to replicate consistently tender meat. As a result, a plethora of research has been conducted to moderate this concern. The University of Nebraska-Lincoln has investigated the impact of different feedstuffs on meat quality with staggering results. Chao (2015), Hart et al. (2019) and Ribeiro et al. (2019) all exhibited inclusions of distillers grains as a model for increased tenderness early postmortem, compared to standard corn rations. This may be a result of distillers grain inclusions impart greater deposition of unsaturated fatty acids within muscle organelles, increasing susceptibility to oxidative mechanisms both ante- and postmortem (Chao et al., 2018). These oxidative mechanisms can alter membrane integrity of organelles, promoting rapid available intracellular molecules for tenderization (Chao et al., 2018).

Therefore, it is conceivable that prolonged oxidative stress prior to death can signal the interaction of oxidative/apoptotic pathways *in vivo* and increase enzymatic activity early postmortem. It is not fully understood how the degree of oxidative stress may impact factors related to *in vivo* myocytes, nor is there a definite understanding on its role with muscle tenderization and color stability in postmortem muscle. As a result,

this study identifies and evaluates mechanisms related to skeletal muscle, using known levels of an oxidative stress promoter prior to harvest on meat quality attributes.

Conclusion

This literature review supports the hypothesis that oxidative stress can manipulate organelle function, promoting ROS-mediated disruption of membrane stability in mitochondria and sarcoplasmic reticulum, upregulating apoptotic pathways necessary during anoxia of postmortem meat, facilitating greater proteolytic potential and muscle tenderization. The exploration of this hypothesis will allow us to use LPS as a mediator of oxidative stress, as *in vivo* oxidative stress is a more viable model system for postmortem tenderization of meat.

Researching this hypothesis will provide valuable insight to the meat industry. If there is a link between proteolytic degradation and the manipulation of organelle confirmation prior to slaughter which promotes tenderness, it could be possible to use oxidative stress as an indicator to enhance meat tenderness, especially when investigating management strategies which influence oxidative stress, such as diet, genetic selection, and environmental control. Additionally, in dealing with muscle cells, there is great potential to understand the biochemical behavior of oxidative stress, in particular how varying degrees of oxidative stress can impact ROS/RNS interrelationships with antioxidant response mechanisms and organelle function. The parabolic nature of oxidative stress is yet to be understood, and there is much knowledge to gain in this biochemical phenomenon.

MATERIALS AND METHODS

Manuscript – Oxidative Stress as a measure for Postmortem Meat Quality

Lambs

A total of 29 cross-bred wethers (initial BW = 29 ± 2.68 kg) were group housed (University of Nebraska Life Sciences Annex in Lincoln, NE) for 42 days on a standard finishing ration (Appendix I) developed at Eastern Nebraska Research and Extension Center (ENREC; Mead, NE). After acclimation period, lambs were blocked and stratified by BW prior to being randomly assigned to a treatment. Lambs were individually fed according to BW to maintain a 0.34kg/day weight gain for the remainder of the in vivo analysis.

Lipopolysaccharide treatments

Lambs were randomly assigned a 2 mL intravenous injection of either saline control (Control), 50ng LPS/kg BW (LPS50) or 100ng LPS/kg BW (LPS100) treatment group. Concentrations of lipopolysaccharide were determined based off of previous research performed at University of Nebraska-Lincoln using sheep as a model for LPS injections. Three injections would be administered across a 12-day immune challenge, with a subsequent injection occurring after every 72 h period. Injection times were partitioned across the 12-day timeline in two groups with 7-8 lambs being administered in each group. Two immune challenge cycles were completed with lambs blocked by BW to determine which cycle they would participate, allowing all animals to reach consistent final BW prior to harvest. All methods of live animal handling were approved by UNL Institutional Animal Care and Use Committee (IACUC).

Rectal temperatures and respiration rates were taken at time 0, 1, 2, 4, 8, 12, 24, 48, and 72 h post-injection time (0600 h). Lambs were secured on a trim stand prior to taking readings. Rectal temperature was completed using a Vicks® Thermometer (ComfortFlex®, V966US, Marlborough, MA) after being inserted 2.54cm from the tip of the thermometer into the rectum of the lamb and held for 10 s. Respiration rate was taken by evaluating the number of exhales taken in a 15 s span. Respiration rates were then calculated up to total breaths per min.

Sample collection

Upon completion of the final 72 h cycle, lambs were held 48 h prior to slaughter, then transported to the Loeffel Meat Lab (Lincoln, NE). Within 30 min of exsanguination, an 80 g pre-rigor sample of posterior Longissimus lumborum from CON and HIGH treatment groups were extracted using a scalpel blade (Feather Safety Razor Co. LTD., No. 11 2976, Osaka, Japan). Pre-rigor sample was denuded of exterior fat and connective tissue, then placed in a 2 mL cryotube (Cryogenic Vial CryoClear 3012 Globe Scientific, Mahwah, N.J.) and frozen for future analysis (-80°C). The carcasses were tagged with a university number that corresponded to the animal and lab ID, then chilled for 24 h.

Fabrication

After 24 h postmortem, sample primals were split between the 8th and 9th rib extending to the last lumber vertebrae. Primals were split down the spine using a band-saw (Biro MFG. Co., Model 3334, Marblehead, OH), and each side was randomly

assigned to one of two aging periods (1 or 14 d). Longissimus muscle was deboned and removed of excess subcutaneous fat. Four 2.54 cm chops were fabricated adjacent to one another at each aging period (one to measure lipid and protein oxidation for 0 d of RD, and three to measure Warner-Bratzler shear force for 0 d retail display). One 2.54 cm chop at each aging period was trimmed of all subcutaneous fat, and utilized to measure visual discoloration, objective color, lipid oxidation, and protein oxidation for 7 d of RD. One 2.54 cm chop at each aging period was cut and utilized for analysis of pH and proximate composition. A 5.08 cm chop was cut the at the most posterior point of the primal and utilized for subsequent laboratory analysis. An additional 2.54 cm chop was cut at the most anterior point of the primal and was used as extra laboratory analysis sample. At d 1, chops for laboratory analysis, pH and proximate composition, extra laboratory analysis were vacuum packaged (MULTIVAC 500, Multivac, Inc., Kansas City, MO) in Prime Source Vacuum pouches (3 mil STD barrier, Prime Sources, St. Louis, MO). Steaks for laboratory analysis, pH and proximate composition were frozen for further analysis (-80°C). The remaining halved portion of loins lined with Boneguard (Boneguard Traditional Perforated, JVR Industries, Lancaster, NY) and aged (2°C) under dark storage. All chops were separated from the loin starting from the anterior end of the loin. The same fabrication map (Appendix II) was used for all aging periods. At all aging periods, samples for color, lipid, and protein oxidation analysis were placed on foam trays (21.6 x 15.9 x 2.1 cm, Styro-Tech, Denver, CO) and overwrapped with an oxygen permeable film (Prime Source PSM 18 #75003815, Bunzl Processors Division, North Kansas City, MO) Trays were placed under simulated RD conditions for 7 d (3°C under white fluorescence lighting at 1000 to 1800 lux) and randomly rotated daily. All frozen

steaks utilized for laboratory analysis and lipid/ protein oxidation were tempered enough to finely dice, freeze in liquid nitrogen and then powdered in a metal cup blender (Model 51BL32, Waring Commercial, Torrington, CT) on May 13/14 and July 26, 2019 and held for 11d and 23d respectively, at -80°C until further analysis.

RNA transcriptomics

Total RNA extraction from muscle was completed using RNeasy Fibrous Tissue Mini Kit (QIAGEN, #74704, Hilden, Germany) and RNase-Free DNase Set (QIAGEN, #79254, Hilden, Germany). Completely sterilize work station with 100% ethanol solution prior to start of RNA extraction. 30 mg of pre-rigor muscle was diced and added to a 1.5 mL conical tube, added with 300µL of RLT solution, and vortexed until broken up.

Homogenize with hand-held pestle for 30 s. Add 10µL proteinase K solution and 590µL RNase-free water and mix thoroughly by pipetting. Incubate at 55°C for 10 min then centrifuge for 3 min at 10,000 x g. Transfer supernatant (approximately 900 µL) into a new 2 mL microcentrifuge (2mL safe-lock tube; 02236352, Eppendorf AG, Hamburg, Germany), avoiding transfer of any pellet. Add 450 µL of 100% ethanol to the cleared lysate. Mix well by pipetting up and down. Transfer 700 µL of the sample to a RNeasy Mini spin column placed in a 2 mL collection tube. Centrifuge for 15 s at 8,000 x g, discard the flow-through. Add 350 µL Buffer RW1 to the RNeasy spin column.

Centrifuge for 15 s at 8,000 x g to wash the membrane, discard the flow-through. Add 10µL DNase I stock solution to 70 µL Buffer RDD. Mix by gently inverting the tube, and centrifuge briefly to collect residual liquid from the sides of the tube. Add the DNase I incubation mix (80 µL) directly to the RNeasy spin column membrane, and place on the

benchtop for 15 min. Add 350 μ L Buffer RW1 to the RNeasy spin column. Centrifuge for 15 s at 8,000 x g. Discard the flow-through. Add 500 μ L Buffer RPE to the RNeasy spin column. Centrifuge for 15 s at 8,000 x g. Discard the flow-through. Add 500 μ L Buffer RPE to the RNeasy spin column. Centrifuge for 2 min at 8,000 x g. Place the RNeasy spin column in a new 2 mL collection tube, and discard the old collection tube with the flow-through. Centrifuge at 14,000 x g for 1 min. Place the RNeasy spin column in a new 1.5 mL collection tube. Add 30-50 μ L RNase-free water directly to the RNeasy spin column membrane. To elute the RNA, centrifuge for 1 min at 8,000 x g. For RNA amplification, transfer samples to University of Nebraska-Medical Center (Next Generation Sequencing, Genome Core Facility, Omaha, NE, USA).

Warner-Bratzler shear force

Three chops (2.54 cm) were measured for tenderness via Warner-Bratzler Shear Force (WBSF) per sample. Internal temperature and weight were measured prior to cooking using a quick disconnect T-type thermocouple (TMQSS-062U-6, OMEGA Engineering, Inc., Stamford, CT) and a handheld thermometer (OMEGA 450-ATT, OMEGA Engineering, Inc., Stamford, CT) in geometric center of steaks. All chops were cooked to an internal temperature of 35°C and turned over until they reached a target temperature of 70°C on an electric indoor grill (Hamilton Beach-31605A, Hamilton Beach Brands, Glen Allen, VA). After cooking, final weights were recorded. The steak was then bagged (PB-90-C, .85 mil., 6x3x15in.) and stored overnight at 2°C. The following day, 2 (1.27 cm diameter) cores per chop were removed with a drill press parallel to muscle fibers and sheared using a Food Texture Analyzer (TMS-Pro, Food

Technology Corp., Sterling, VA.) with a triangular Warner-Bratzler blade. The mean of 6 cores was calculated for statistical analysis.

Objective color and subjective color:

Objective color measurements were taken once daily for 7 days during simulated retail display at all aging time points. Chops (2.54 cm) were placed on Styrofoam trays (21.6 x 15.9 x 2.1 cm, Styro-Tech, Denver, CO), overwrapped with oxygen permeable film (Prime Source PSM 18 #75003815, Bunzl Processors Division, North Kansas City, MO), and placed under retail display conditions (3°C under white fluorescence lightening at 1,000 to 1,800 lux). Commission international de l'éclairage (CIE) L* a* b* values were obtained using a Minolta CR-400 colorimeter (Minolta, Osaka, Japan) set with a D65 illuminant, 2°C, with an 8 mm diameter measurement area. Three measurements were made per chop and the mean was calculated for statistical analysis. The colorimeter was calibrated daily with a white ceramic tile (Calibration Plate, Serial No. 14933058, Konica Minolta, Japan). Lightness (L*) is measured with a range from 0 (black) to 100 (white), a* measures redness with the range between red (positive) and green (negative), and b* is a measure of yellowness from yellow (positive) to blue (negative). Color readings were recorded at the same time each day.

Visual discoloration was assessed daily during the 7 d of RD utilizing 5 trained panelists comprised of graduate students from the University of Nebraska. Panelists were trained using a standardized discoloration guide (Appendix VI). Discoloration % was approximated from 0% to 100% with 0% meaning no discoloration present and 100%

being a fully discolored chop. Chops were randomly rotated daily to minimize location effects.

Proximate composition

Moisture, fat, and ash (%) of raw meat samples were determined. Samples were measured in triplicate in Whatman #2 paper filter paper and fat was extracted with anhydrous ether. After identifying and recording weights of the folded filter papers with corresponding paper clips to hold packets closed, these were tared out and powdered meat samples (2 g) were weighed onto the filter paper and then closed with the paper clip. Samples were then placed in a Soxhlet tubes and the boiling flasks were filled with 400 mL of ether. Once in place, water was opened to enter the condensers and each individual burner was turned on. After 48 h, burners were turned off and allowed to cool completely. Samples were air-dried under a fume hood for 2 h, then placed in a drying oven (105°C) overnight prior to recording final dry weight. In order to calculate final fat percentage, the final equation was used: % Fat = ((Pre-extraction wet weight with filter paper and paper clip – Post-extraction dry sample weight)/sample weight) * 100) - % Moisture. Moisture and ash (%) were calculated with a LECO Thermogravimetric Analyzer in duplicate (Model 604-100-400, LECO Corporation, St. Joseph, MI), see Appendix IV for information. Moisture was determined in nitrogen atmosphere with a start temperature of 25°C and an end temperature of 130°C (17 min ramp rate). Ash was determined in oxygen atmosphere with a start temperature of 130°C and an end temperature of 600°C (30 min ramp rate). Protein was determined by difference.

Sarcomere length

Sarcomere length was determined using the helium-neon laser diffraction method described by Cross et al. (1981) and Dolazza and Lorenzen (2014). A few flecks of powdered meat sample were placed on a clear glass microscopic slide. A single drop of 0.25M sucrose solution was added to the slide and topped with a glass coverslip. The distance to the top of the slide from the base of the laser was 100mm. A sheet of paper was placed below the stand in order to mark the two diffraction bands. Six sarcomeres per sample were determined and sarcomere length (μm) was determined by the equation provided by Cross et al. (1981):

Sarcomere length (μm) =

$$\mu = \frac{0.6328 \times D \sqrt{\left(\frac{T^2}{D}\right) + 1}}{T}$$

Where:

0.6328 = 632.8 (the wavelength of the laser) $\times 10^{-3}$

D = distance from specimen to diffraction pattern screen (100mm)

T = spacing between diffraction bands (mm)

pH analysis

Powdered sample from chops from all aging periods with 0 d RD were weighed out in 5 g duplicates into 250 mL plastic beakers and placed on a stir plate. Forty-five mL of distilled deionized water and a magnetic stir bar were added to ensure constant mixing during the measurement process. The pH was measured using a pH meter (Orion

410Aplus: ThermoFisher Scientific; Waltham, MA) that was calibrated using 4.0, 7.0, 10.0 standards. The mean measurement of the duplicates was utilized for all analysis.

Fatty acids

Fatty acid profiles were obtained via gas chromatography as described by (Folch et al., 1987). After extraction, lipids were converted to fatty acid methyl esters according to Morrison and Smith (1964) and Metcalfe et al. (1966). One g of powdered sample was homogenized with 5 mL of 2:1 chloroform: methanol and allowed to sit at room temperature (23°C) for 1 h. After, samples were filtered through Whatman #2 paper, brought up to a final volume of 10mL with 2:1 chloroform: methanol, and vortexed for 5 s with 2mL of 0.74% KCl. Samples were centrifuged (1,000 x g for 5 min at 5°C) and the top layer was aspirated off. After centrifugation, samples were dried on a heating block at 60°C under nitrogen purge. Once dry, 1 mL of 0.5 M NaOH in methanol was added, vortexed (5 s), and again heated at 100°C for 10 min. One mL of 14% Boron Trifluoride in methanol was added, vortexed (5 s), and again heated at 100°C, this time for 5 min. Two mL of saturated salt solution and 2 mL of hexane was added and vortexed (5 s). Samples were then centrifuged (1,000 x g for 5 min at 5°C) and the hexane layer removed and analyzed using gas chromatography (TRACE 1310 Gas Chromatograph; ThermoFisher Scientific, Waltham, MA). Fatty acids were separated using a Chrompack CP-Sil 88 capillary column (0.25mm by 100mm; Inlet temp: 260°C, Oven: 140°C hold for 5 min, increase at 4°C/min to 240°C and hold for 15 min. FID temp: 280°C. Injected at 30:1 ratio) and identified based on their retention times compared to known commercial standards (NU-Check Prep, Inc., Elysian, MN; #GLC-68D, GLC-79, GLC-

87, GLC-455, and GLC-458). The percentage of fatty acids were determined by the peak areas in the chromatograph and values were converted to mg/100g tissue: Fatty acid mg/100g tissue = (% of fatty acid peak area * fat content of samples) * 100.

Free calcium concentration

Free calcium was quantified according to the procedure described by Parrish et al. (1981) with modifications. Three grams of powdered sample were centrifuged at 196,000 x g (Beckman Optima XPN-90 Ultracentrifuge, Type 50.2 Ti rotor, Beckman Coulter, Brea, CA) at 4°C for 30 min. Seven hundred microliters of the supernatant were collected and treated with 0.1 mL of 27.5 trichloroacetic acid (TCA). Samples were centrifuged at 6,000 x g (accuSpin Micro 17R, ThermoFisher Scientific, Waltham, MA) for 10 min at 4°C. Four hundred µL of supernatant were transferred to a syringe, and the volume was brought to 4 mL with deionized, distilled water. The diluted sample was filtered through a 13 mm diameter Millex-LG 0.20 µm syringe filter (Millipore, Bedford, MA). Calcium concentration was quantified at Ward Laboratories (Kearney, NE) using an inductively-coupled plasma emission spectrometer (iCAP 6500 Radial; Thermo Electron, Cambridge, UK) with an appropriate calcium concentration standard.

Lipid oxidation (TBARS)

Lipid oxidation was determined using thiobarbaturic acid reactive substances values (TBARS) for all aging periods at 0 and 7 d RD described (Ahn et al., 1998). Five grams of powdered meat were placed into a 50 mL conical tube to which 14 mL of distilled deionized water were added and 1 mL of butylated hydroxyanisole (BHA)

solution (10% BHA: 90% ethanol). Samples were homogenized using a Polytron (POLYTRON® Kinimatica CH-6010, Switzerland) for 15 s at medium to high speed. The samples were centrifuged (2,000 x g for 5 min at 10°C) and one mL of supernatant was transferred into a 15 mL conical tube with 2 mL of 2,4,6-tribromoanisole (TBA) 2,4,6-trichloroanisole (TCA) solution (15% and 20 mM TBA in deionized distilled water). Tubes were then placed in a 70°C water bath for 30 min. After 30 min, tubes were cooled for at least 10 min in a water bath (22°C) and centrifuged (2,000 x g for 15 min at 10°C). Two hundred µL of supernatant was transferred to a 96-well plate in duplicate (Microtest III sterile 96 well flat-bottomed microplate; Becton Dickinson & Company, Lincoln Park, NJ). Absorbance values were then read at 540 nm using a microplate spectrophotometer and compared to known standards (Model Epoch Biotek, Winooski, VT). Results were expressed in mg of malonaldehyde per kg of tissue.

Troponin T

Troponin T degradation was quantified according the procedure described by Chao et al. (2018) with modifications. 3 grams of powdered meat was homogenized with 15 mL of ice-cold rigor buffer (0.1M KCl, 2mM MgCl₂, 1mM EDTA, and 10mM K₂HPO₄; pH 7.4) using a polytron (POLYTRON Kinimatica CH-6010, Switzerland) at medium speed for 5 s bursts for 30 s. Homogenate was filtered through a double-layered cheese cloth and 1.4 mL of homogenate was extracted into an eppendorf tube (2mL safe-lock tube; 02236352, Eppendorf AG, Hamburg, Germany). Tubes were centrifuged at 4,000 x g for 5 min. Supernatant was decanted and pellet was resuspended in 1 mL of rigor buffer. Centrifuge sample at 4,000 x g for 5 min two more times. Decant

supernatant and suspend pellet in 1 mL suspension buffer (0.1M Tris-Base, 1.25mM EDTA, 5% SDS; pH 8) and centrifuged at 4,000 x g for 5 min. Protein concentration was determined using a Pierce bicinchoninic acid protein assay kit (Pierce Biotechnology, Rockford, IL, USA). All samples were diluted to 2 mg/mL with deionized-distilled water and 2x Laemmli buffer (65.8mM Tris-HCl, 2.1% SDS, 26.3% glycerol, 0.01% bromophenol blue) with 2% betamercaptoethanol (1:50) and put on a heating block at 95°C for 5 min. 20µL of sample were loaded on 4-20% Mini-PROTEAN TGX™ precast polyacrylamide gels (Bio-Rad Laboratories, Hercules, CA, USA) with a 10µL pre-stained standard (Precision Plus Protein Kaleidoscope, #1610375, Bio Rad, Hercules, CA) using a Bio-Rad Mini-PROTEIN 2 Cells system (Bio-Rad Laboratories). The system was run at constant voltage of 200V for 60 min with an electrophoresis buffer (1xTris/Glycine/SDS, #161-0732, Bio-Rad Laboratories, Hercules, CA, USA). Proteins in the gels were blotted to polyvinylidene difluoride membranes (0.45µm, Immobion-FL transfer membrane; Millipore) using a Bio-Rad Mini-Trans-Blot Electrophoretic transfer cell (Bio-Rad Laboratories) for 60 min at a constant amperage of 180mA with ice-cold transfer buffer (25mM Tris-base, 192mM Glycine, 20% methanol; pH at 9.2). Membranes were blocked for 2 h in Odyssey Blocking Buffer (LI-COR, Lincoln, NE, USA) and incubated for 60 min at room temperature in monoclonal anti-troponin-T antibody (JLT-12; Sigma-Aldrich, St. Louis, MO, USA) at a dilution of 1:10,000 in Odyssey blocking buffer containing 0.2% TWEEN-20 and 5% non-fat dry milk. Membranes incubated overnight at 4°C, then washed three time with Tris Buffered Saline containing 0.2% TWEEN-20 (TBST) for 10 min and incubated in IRDye 680 LT Conjugated Goat Anti-Mouse IgG1 secondary antibody (LI-COR) at a dilution of 1: 10,000 in Odyssey blocking buffer

containing 0.2% TWEEN-20 for 60 min. Membranes were washed three times with TBST and scanned using Odyssey Infrared Imaging system (LI-COR) at 700nm. Degradation was evaluated by quantifying band intensities (k. pixels) using Odyssey application software version 1.1. Bands ranging from 38 and 35kDa were designated as intact and bands ranging from 30 to 28kDa were designated as degraded. Percent degradation was calculated by $(\text{intensity of degraded bands}/\text{intensity of intact bands}) \times 100$

Desmin

Three grams of powdered meat was homogenized with 15 mL of whole muscle solubilization buffer (2%wt/vol SDS, 10mM sodium phosphate buffer; pH 7.0) using a polytron (POLYTRON Kinimatica CH-6010, Switzerland) at medium speed for 5 s bursts for 30 s. Homogenate was filtered through a double-layered cheese cloth and 1.4 mL of homogenate was extracted into an eppendorf tube (2mL safe-lock tube; 02236352, Eppendorf AG, Hamburg, Germany). Tubes were centrifuged at 1,500 x g for 15 min at 25°C to remove traces of insoluble components. 1 mL of supernatant was collected and into a 1.5 mL eppendorf tube. Protein concentration was determined using a Pierce bicinchoninic acid protein assay kit (Pierce Biotechnology, Rockford, IL, USA). All samples were diluted to 6.4 mg/mL using whole muscle solubilizing buffer. Samples were diluted to a final concentration of 4mg/mL with 50% of gel buffer (3mM EDTA, 3% wt/vol SDS, 30% vol/vol glycerol, 0.001% wt/vol pyronin Y, and 30mM Tris-HCl; pH 8.0) and 10% of betamercaptoethanol then heated on a block for 15 min at 50°C. 30µL of sample were loaded on 4-20% Mini-PROTEAN TGX™ precast polyacrylamide gels (Bio-Rad Laboratories, Hercules, CA, USA) with a 10µL pre-stained standard

(Precision Plus Protein Kaleidoscope, #1610375, Bio Rad, Hercules, CA) using a Bio-Rad Mini-PROTEIN 2 Cells system (Bio-Rad Laboratories). The system was run at constant voltage of 360V for 45 min with an electrophoresis buffer (1xTris/Glycine/SDS, #161-0732, Bio-Rad Laboratories, Hercules, CA, USA). Proteins in the gels were blotted to polyvinylidene difluoride membranes (0.45 μ m, Immobion-FL transfer membrane; Millipore) using a Bio-Rad Mini-Trans-Blot Electrophoretic transfer cell (Bio-Rad Laboratories) for 90 min at a constant voltage of 90V with ice-cold transfer buffer (25mM Tris-base, 192mM Glycine, 20% methanol; pH at 9.2). Membranes were blocked for 60 min in Odyssey Blocking Buffer (LI-COR, Lincoln, NE, USA) and incubated for 60 min at room temperature in monoclonal anti-desmin antibody (DE-U-10 Sigma-Aldrich, St. Louis, MO, USA) at a dilution of 1:10,000 in Odyssey blocking buffer containing 0.2% TWEEN-20 and 5% non-fat dry milk. Membranes were incubated overnight at 4°C, then washed three times with Tris Buffered Saline containing 0.2% TWEEN-20 (TBST) for 10 min and incubated in IRDye 680 LT Conjugated Goat Anti-Mouse IgG1 secondary antibody (LI-COR) at a dilution of 1: 10,000 in Odyssey blocking buffer containing 0.2% TWEEN-20 for 60 min. Membranes were washed three times with TBST and scanned using Odyssey Infrared Imaging system (LI-COR) at 700nm. Degradation was evaluated by quantifying band intensities (k. pixels) using Odyssey application software version 1.1. Bands at 55kDa were designated as intact and bands at 38kDa were designated as degraded. Percent degradation was calculated by $(\text{intensity of degraded bands}/\text{intensity of intact bands}) * 100$.

Isoprostanes

All analysis was completed using OxiSelect™ 8-iso-Prostaglandin F2alpha ELISA Kit (Cell BioLabs, INC., STA-337, San Diego, CA). Weigh out 30mg of powdered meat in a 5 mL eppendorf tube (). Homogenize sample in 2 mL of 2N NaOH using Micropolytron(Pro-Scientific200, Bio-Gen, Oxford, CT) and spin for 20 s, cleaning micropolytron between each sample. Heat homogenized sample in water bath (Thelco 261, GCA Precision Scientific, Chicago, IL) at 45°C for 2 h to ensure hydrolysis. Cool tissue samples to room temperature (~20 min), then neutralize pH using 2 mL of 2N HCl, vortex for 20 s. Split into two 2 mL eppendorf tubes (2mL safe-lock tube; 02236352, Eppendorf AG, Hamburg, Germany). Using microcentrifuge, spin samples at 10,000 x g for 15 min at 4°C. Add 100µL of sample to 100µL of neutralization solution in a 0.6 mL eppendorf tube. Dilute Anti-8-iso-PGF2α Antibody (1:1000) with sample diluent. Add 100µL of the diluted antibody to the Goat Anti-Rabbit Antibody coated plate. Incubate on an oscillation shake at 25°C for 1 h. During incubation, prepare 100 mL of 1x wash buffer by diluting 10x wash buffer concentrate with deionized water. Prepare Isoprostanes standards as labeled in Appendix XIX. After incubation, remove the antibody solution from the wells. Wash wells 5 times with 300µL 1x wash buffer per well. After the last wash, empty the wells and tap microwell plate on absorbent pad or paper towel to remove excess wash solution. Immediately before use, dilute the conjugate (1:80) with sample diluent. Combine 55µL of the 8-iso-PGF2α standard or hydrolyzed sample and 55µL of 8-iso-PGF2alpha-HRP conjugate in a microtube and mix thoroughly. Transfer 100µL of the combined solution per well. A well containing sample diluent can be used as a blank. Incubate plate for 1 h at 25°C on an orbital shaker. After incubation,

remove the combined solution from the wells. Wash 5 times with 300 μ L of 1x wash buffer per well. After the last wash, empty wells and tap microwell plate on absorbent pad or paper. Add 100 μ L of substrate solution to each well and incubate at 25°C for 30 min on oscillate shaker. After incubation, add 100 μ L of stop solution to each well. Results should be read immediately on a microplate reader using 450nm as the target wavelength. Units of isoprostane content designated as picogram/mL.

Statistical analysis

Statistical analysis was conducted with SAS (version 9.4, Cary, NC). Objective and subjective color data were analyzed as a split-plot repeated measures design with treatment as the whole-plot, aging period as the split-plot and retail display as the repeated measures. Tenderness, troponin t, desmin, calcium, and pH were analyzed as a split-plot design with treatment as the whole-plot and aging period as the split-plot. Lipid oxidation, free thiols and carbonyls were a split-split-plot design with treatment as the whole plot, aging period as the split-plot and retail display time as the split-split-plot. Transcriptomics, sarcomere length, fatty acids, cytochrome c, and isoprostanes were analyzed as a completely randomized design. Data were analyzed using the PROC GLIMMIX procedure of SAS and animal was the experimental unit. All means were separated with the LS MEANS statement with an α level of 0.05 and tendencies were considered at an α level of 0.15.

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Oxidative Stress as a measure of Postmortem Meat Quality

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Abstract

The objective of this study was to evaluate the effects of different levels of lipopolysaccharide (LPS)-mediated oxidative stress on fresh meat quality. Crossbred lambs ($n = 29$) were blocked by weight and fed a standard finishing ration for the duration of the study. Lambs were individually housed, and treatment groups were administered one of three injections every 72 hours across a three-injection (9-day) cycle: a saline control (**Control**), 50 ng LPS/kg bodyweight (**LPS50**), or 100 ng LPS/kg bodyweight (**LPS100**). Rectal temperatures were measured to indicate inflammatory response. Lambs were harvested at the Loeffel Meat Laboratory, and 80 mg of pre-rigor *Longissimus lumborum* were collected in Control and LPS100 treatments within thirty minutes postmortem for RNA analysis. Loins were split and randomly assigned for 1 or 14 d of aging. Chops were fabricated after aging and placed under retail display (**RD**) conditions for 0 or 7 d. Animal was the experimental unit. Lipopolysaccharide-treated lambs had increased ($P < 0.05$) rectal temperatures at 1, 2, 4, and 24 h post-injection. Transcriptomics exhibited significant ($P_{\text{raw}} < 0.05$) upregulation in RNA pathways related to generation of oxidative stress in LPS100 compared to Control. A trend was found for tenderness (Warner-Bratzler shear) ($P = 0.10$), with chops from LPS50 having a lower shear force compared with Control at 1 d postmortem. The LPS50 treatment exhibited greater troponin T degradation ($P = 0.02$) compared to all treatments at 1 d. Aging decreased WBSF ($P < 0.0001$), increased free calcium concentration ($P < 0.0001$), pH ($P < 0.0001$), and proteolytic degradation ($P < 0.0001$) across all treatments. After 7 d of RD, following aging periods, chops increased discoloration as RD increased ($P <$

0.0001), with Control chops aged for 14 d being the most discolored. Chops from lambs given LPS had higher ($P < 0.05$) a^* values compared to control at 14 d of aging. The L^* values were greater ($P < 0.05$) in LPS100 compared to both LPS50 and Control. Aging tended ($P = 0.0608$) to increased lipid oxidation (TBARS) during RD across either aging period. There were no significant differences ($P > 0.05$) in sarcomere length, proximate composition, fatty acid composition, or isoprostane content. These results suggest that defined upregulation of oxidative stress has no detriment on fresh meat color, but may alter biological pathways responsible for muscle composition and enzymatic processes, resulting in changes in tenderness early postmortem.

Key words: apoptosis, color stability, lamb, meat quality, oxidative stress, tenderization

Introduction

During postmortem aging, the tenderization of muscle relies on several biological mechanisms, including endogenous enzymes, such as calpains, caspases, and proteasomes (Ouali et al., 2006; Bhat et al., 2018). These endogenous mechanisms activate early postmortem with the utilization of free calcium released from mitochondria and sarcoplasmic reticulum (Rossi and Dirksen, 2006; Santo-Domingo and Demareux, 2010). With regards to postmortem aging, an increasing number of investigations have focused on the influence of apoptosis on meat quality. Kemp and Parr (2012) indicated apoptotic events can contribute to proteolytic degradation of many structural proteins during muscle aging.

Apoptosis is the mechanism responsible for regulated cell death, portrayed by multiple biochemical and molecular pathways (Sierra and Olivan, 2013). This intrinsic

mechanism involves mitochondria, as increased apoptotic events promote mitophagy, as indicated by the release of cytochrome c from the mitochondrial membrane into the cytoplasm (Ott et al., 2007; Kagan et al., 2009; Ke et al., 2017). Cytosolic cytochrome c binds to Apaf-1 genes, responsible for initiating enzymatic proteins, to produce apoptosomes, large quaternary proteins used to initiate caspase systems and begin proteolytic events (Porter and Jänicke, 1999; Momeni, 2011). Initiation of apoptotic activity has been linked to oxidative stress-mediated events (Slimen et al., 2014).

The onset of oxidative stress results from the overwhelming production of reactive oxygen species (ROS) compared to homeostatic endogenous antioxidants present within the system (Powers et al., 2011). As the name implies, ROS are highly reactive substances primarily produced as by-products during oxidative phosphorylation (Paradies et al., 2001). Additionally, ROS act as signaling molecules to upregulate a homeostatic response (Dröge, 2002). This action facilitates antioxidant enzymes (superoxide dismutase, catalase, glutathione) capable of changing ROS to more stable products within muscle cells. However, conditions of oxidative stress overwhelm antioxidant function, allowing ROS to alter protein, lipid, or nucleic acid morphology and functionality (Bekhit et al., 2013). Concurrently, ROS can interact with nitrosative species (nitric oxide), producing hybrid radicals which can target proteins responsible for organelle function (Stamler and Meissner, 2001). Given the right conditions, oxidative stress factors have been theorized to impact muscle cells and create conditions critical to meat quality.

Intrinsic and extrinsic conditions promoting oxidative stress have been implicated to alter meat quality (Warner et al., 2005; Ponnampalam et al., 2017; Wang et al., 2018; Mitacek et al., 2019). However, there is inconsistency within the literature describing the

impact of oxidative stress factors on meat quality, as this phenomenon is theorized to be dependent on a multitude of factors. These factors include *degree* and duration of oxidative stress, source of oxidative stress generation (*in vivo* vs *in vitro*), the influence of individual oxidative and nitrosative species on cellular constituents, and composition of muscle tissue (Cottrell et al., 2015; Niu et al., 2016; Ke et al., 2017; Wang et al., 2018). Concurrently, there is a gap in the literature relating specific genetic pathways and their oxidative stress-mediated regulation on the impact of meat quality.

The mechanism of meat tenderization is well-recognized. However, the impact of oxidative stress on skeletal muscle, and its influence on factors critical to meat quality has yet to be understood. Additionally, the impact of oxidative stress and its relationship towards the muscle transcriptome are not fully understood. Therefore, the objective of this study was to evaluate the effects of controlled oxidative stress *in vivo* on oxidative biomarkers within the muscle transcriptome and meat quality attributes, including tenderness, color stability, and lipid oxidation.

Materials and Methods

All animal use protocols were approved by the University of Nebraska-Lincoln's Institutional Animal Care and Use Committee [Protocol No. 1751].

Lambs

A total of 29 cross-bred (Hampshire x Dorset x Polypay) wethers (initial BW = 29 ± 2.68 kg) were group housed (University of Nebraska Life Sciences Annex in Lincoln, NE) for 42 days on a standard finishing ration (Appendix I) developed at Eastern

Nebraska Research and Extension Center (ENREC; Mead, NE). After the acclimation period, lambs were blocked and stratified by BW prior to being randomly assigned to a treatment. Lambs were individually fed according to BW to maintain a 0.34kg/day weight gain for the remainder of the in vivo analysis.

Lipopolysaccharide treatments

Lambs were randomly assigned a 2 mL intravenous injection of either saline control (Control), 50ng LPS/kg BW (LPS50) or 100ng LPS/kg BW (LPS100) treatment group. Concentrations of lipopolysaccharide O111: B4 (L2630; Sigma-Aldrich) were determined based on previous research performed at University of Nebraska-Lincoln using sheep as a model for LPS injections. Three injections were administered across a 12-day immune challenge, with a subsequent injection occurring after every 72 h period. Injection times were partitioned across the 12-day immune challenge, with two sets of 14-15 lambs being administered in each group. Two immune challenge groups were completed with lambs blocked by BW to determine which cycle they would participate, allowing all animals to reach consistent final BW prior to harvest. All methods of live animal handling were approved by UNL Institutional Animal Care and Use Committee (IACUC).

Rectal temperatures

Rectal temperatures and respiration rates were measured at time 0, 1, 2, 4, 8, 12, 24, 48, and 72 h post-injection time (0600 h). Lambs were secured on a trim stand prior to taking readings. Rectal temperature was completed using a Vicks® Thermometer

(ComfortFlex®, V966US, Marlborough, MA) by inserting 2.54cm in from the tip of the anus to the rectum of the lamb and held for 10 s. Temperatures were recorded as degrees Centigrade.

Muscle Sample collection

Upon completion of the final 72 h cycle, lambs were held 48 h prior to slaughter, then transported to the Loeffel Meat Lab (Lincoln, NE). Within 30 min of exsanguination, an 80 mg pre-rigor sample taken from the posterior end of the Longissimus lumborum from Control and LPS100 treatment groups. Samples were extracted using a scalpel blade sterilized in ethanol in between sampling (Feather Safety Razor Co. LTD., No. 11 2976, Osaka, Japan). Pre-rigor samples were removed of exterior fat and connective tissue, then placed in a 2 mL cryotube (Cryogenic Vial CryoClear 3012 Globe Scientific, Mahwah, N.J.) and frozen for future analysis (-80°C). The carcasses were tagged with a university number that corresponded to the animal and lab ID, then chilled for 24 h.

Fabrication

After 24 h postmortem, carcasses were fabricated and the loin portion from the 9th rib to the last lumbar vertebrae was retained. Each loin section was split down the spine using a band-saw (Biro MFG. Co., Model 3334, Marblehead, OH). Within each animal, sides were randomly assigned to one of two aging periods (1 or 14 d). Longissimus muscle was deboned and removed of excess subcutaneous fat. Beginning at the posterior end, one 5.08 cm chop was cut and utilized for all laboratory analysis at each aging

period. One 2.54 cm chop at each aging period was utilized for analysis of pH and proximate composition. Three 2.54 cm chops were fabricated adjacent to one another for each aging period and analyzed for tenderness using Warner-Bratzler shear force for 0 d retail display. One 2.54 cm chop was cut to measure lipid and protein oxidation for 0 d of RD. One 2.54 cm chop at each aging period was trimmed of all subcutaneous fat, and utilized to measure visual discoloration, objective color, lipid oxidation, and protein oxidation for 7 d of RD. Lastly, a 2.54 cm chop was cut at the most anterior point of the primal and was used as extra laboratory analysis sample. At d 1 aging, chops for laboratory analysis, pH and proximate composition, extra laboratory analysis were vacuum packaged (MULTIVAC 500, Multivac, Inc., Kansas City, MO) in Prime Source Vacuum pouches (3 mil STD barrier, Prime Sources, St. Louis, MO). Chops for laboratory analysis, pH and proximate composition were frozen for further analysis (-80°C). The remaining halved portion of loins were lined with Boneguard (Boneguard Traditional Perforated, JVR Industries, Lancaster, NY), vacuum packaged, and aged (2°C) under dark storage for 14 days total. All chops were separated from the loin starting from the anterior end of the loin. The same fabrication map (Appendix II) was used for all aging periods. For both aging periods, samples for color, lipid, and protein oxidation analysis were placed on foam trays (21.6 x 15.9 x 2.1 cm, Styro-Tech, Denver, CO) and overwrapped with an oxygen permeable film (Prime Source PSM 18 #75003815, Bunzl Processors Division, North Kansas City, MO) Trays were placed under simulated RD conditions for 7 d (3°C under white fluorescence lighting at 1000 to 1800 lux) and randomly rotated daily. All frozen steaks utilized for laboratory analysis and lipid/ protein oxidation were tempered enough to finely dice, freeze in liquid nitrogen

and then powdered in a metal cup blender (Model 51BL32, Waring Commercial, Torrington, CT) on May 13/14 and July 26, 2019 and held for 11d and 23d respectively, at -80°C until further analysis.

RNA transcriptomics

Total RNA isolation from muscle was completed using the RNeasy Fibrous Tissue Mini Kit (QIAGEN, #74704, Hilden, Germany) and RNase-Free DNase Set (QIAGEN, #79254, Hilden, Germany). Utilize RNaseZap prior to start of RNA isolation to remove RNase from all work stations. Less than 30 mg of frozen pre-rigor muscle was weighed and added to a 1.5 mL microcentrifuge tube. Tissue was vortexed in 300 μ L of RLT solution (3 μ L β -mercaptoethanol mixed with 300 μ L of Buffer RLT) for up to 30 s and then homogenized with a hand-held pestle for another 30 s. 10 μ L of Proteinase K solution and 590 μ L of RNase-free water were mixed thoroughly by pipetting and added to the sample. The sample solution was incubated at 55°C for 10 min then centrifuged for 3 min at 10,000 x g. Approximately 900 μ L of supernatant was transferred into a 2 mL microcentrifuge tube, while avoiding transfer of pellet. Half the supernatant volume in 100% ethanol was added to the cleared lysate and mixed by pipetting. 700 μ L of the sample was transferred to an RNeasy Mini spin column placed in a 2 mL collection tube. Samples were centrifuged for 15 s at 8,000 x g. Flow-through was discarded. The rest of the sample was transferred to the column and re-centrifuged. Three-hundred fifty μ L of Buffer RW1 was added to the spin column and centrifuged for 15 s at 8,000 x g. The flow-through was discarded. 10 μ L of DNase I stock solution (prepared using the handbook instructions) was added to 70 μ L Buffer RDD and mixed by inverting the tube.

DNase I incubation mix (80 μ L) was added directly to the spin column membrane, and incubated at room temperature for 15 min. 350 μ L of Buffer RW1 was added to the spin column and centrifuged for 15 s at 8,000 x g. Filtered solution was discarded. Five-hundred μ L of Buffer RPE was added to the spin column and centrifuged for 15 s at 8,000 x g. Filtered solution was discarded. 500 μ L of Buffer RPE was added to the spin column and centrifuged for 2 min at 8,000 x g. The spin column was placed into a new 2 mL collection tube and centrifuged for 1 min at 14,000 x g. The spin column was placed in a 1.5 mL microcentrifuge tube. 50 μ L of RNase-free water was directly added to the spin column membrane and centrifuged for 1 min at 8,000 x g. Samples were sent to the University of Nebraska-Medical Center (Next Generation Sequencing, Genome Core Facility, Omaha, NE, USA) for poly-A+ library preparation and sequencing (150bp paired-end).

Resulting data were quality trimmed using Trim Galore! , and aligned to the Oar_rambouillet_v1.0 reference genome STAR (Dobin et al., 2016). Differential expression (control vs LPS treatment) was evaluated using transcript counts in DESeq2 (Love et al., 2014). Loci with $P_{adj}<0.05$ were considered to be differentially expressed; those with $P_{raw}<0.05$ were utilized for pathway exploration in Ingenuity Pathway Analysis (Qiagen).

Warner-Bratzler shear force (WBSF)

Three chops (2.54 cm) from each side were measured for tenderness via Warner-Bratzler Shear Force (WBSF) per sample. Internal temperatures were measured prior to cooking using a quick disconnect T-type thermocouple (TMQSS-062U-6, OMEGA

Engineering, Inc., Stamford, CT) with a handheld thermometer (OMEGA 450-ATT, OMEGA Engineering, Inc., Stamford, CT) in geometric center of steaks. Weights of chops were collected prior to cooking using a precision balance scale (PL6001E, Mettler Toledo, Hogentogler and Co. Inc., Columbia, MO). All chops were cooked to an internal temperature of 35°C and turned over until they reached a target temperature of 70°C on an electric indoor grill (Hamilton Beach-31605A, Hamilton Beach Brands, Glen Allen, VA). After cooking, final weights were recorded. The steak was then bagged (PB-90-C, .85 mil., 6x3x15in. PITT PLASTICS, Pittsburg, KS) and stored overnight at 2°C. The following day, 2 (1.27 cm diameter) cores per chop were removed with a drill press parallel to muscle fibers and sheared using a Food Texture Analyzer (TMS-Pro, Food Technology Corp., Sterling, VA.) with a triangular Warner-Bratzler blade. The mean of 6 cores was calculated for statistical analysis.

Troponin T

Troponin T degradation was quantified according to the procedure described by Chao et al. (2018) with modifications. Three grams of powdered meat was homogenized with 15 mL of ice-cold rigor buffer (0.1M KCl, 2mM MgCl₂, 1mM EDTA, and 10mM K₂HPO₄; pH 7.4) using a polytron (POLYTRON Kinimatica CH-6010, Switzerland) at medium speed for 5 s bursts for 30 s. Homogenate was filtered through a double-layered cheese cloth and 1.4 mL of homogenate was placed into an Eppendorf tube (2mL safe-lock tube; 02236352, Eppendorf AG, Hamburg, Germany). Tubes were centrifuged at 4,000 x g for 5 min at 4°C. Supernatant was decanted and pellet was resuspended in 1 mL of rigor buffer. Samples were centrifuged at 4,000 x g for 5 min two additional times.

Supernatant was decanted and the suspended pellet was mixed in 1 mL suspension buffer (0.1M Tris-Base, 1.25mM EDTA, 5% SDS; pH 8) and centrifuged at 4,000 x g for 5 min. Protein concentration was determined using a Pierce bicinchoninic acid protein assay kit (Pierce Biotechnology, Rockford, IL, USA). All samples were diluted to 2 mg protein/mL with deionized-distilled water and 2x Laemmli buffer (65.8mM Tris-HCl, 2.1% SDS, 26.3% glycerol, 0.01% bromophenol blue) with 2% betamercaptoethanol (1:50) and put on a heating block at 95°C for 5 min. 20µL of sample were loaded on 4-20% Mini-PROTEAN TGX™ precast polyacrylamide gels (Bio-Rad Laboratories, Hercules, CA, USA) with a 10µL pre-stained standard (Precision Plus Protein Kaleidoscope, #1610375, Bio Rad, Hercules, CA) using a Bio-Rad Mini-PROTEIN 2 Cells system (Bio-Rad Laboratories). The system was run at constant voltage of 200V for 60 min with an electrophoresis buffer (1xTris/Glycine/SDS, #161-0732, Bio-Rad Laboratories, Hercules, CA, USA). Proteins in the gels were blotted to polyvinylidene difluoride membranes (0.45µm, Immobion-FL transfer membrane; Millipore) using a Bio-Rad Mini-Trans-Blot Electrophoretic transfer cell (Bio-Rad Laboratories) for 60 min at a constant amperage of 180mA with ice-cold transfer buffer (25mM Tris-base, 192mM Glycine, 20% methanol; pH at 9.2). Membranes were blocked for 2 h in Odyssey Blocking Buffer (LI-COR, Lincoln, NE, USA) and incubated for 60 min at room temperature in monoclonal anti-troponin-T antibody (JLT-12; Sigma-Aldrich, St. Louis, MO, USA) at a dilution of 1:10,000 in Odyssey blocking buffer containing 0.2% TWEEN-20 and 5% non-fat dry milk. Membranes were incubated overnight at 4°C and then washed three times with Tris Buffered Saline containing 0.2% TWEEN-20 (TBST) for 10 min. Membranes were incubated in IRDye 680 LT Conjugated Goat Anti-Mouse

IgG1 secondary antibody (LI-COR) at a dilution of 1: 10,000 in Odyssey blocking buffer containing 0.2% TWEEN-20 for 60 min. Membranes were washed three times with TBST and scanned using Odyssey Infrared Imaging system (LI-COR) at 700nm. Degradation was evaluated by quantifying band intensities (k. pixels) using Odyssey application software version 1.1. Bands ranging from 38 and 35kDa were designated as intact and bands ranging from 30 to 28kDa were designated as degraded. Percent degradation was calculated by $(\text{intensity of degraded bands}/\text{intensity of intact bands}) * 100$.

Desmin

Three grams of powdered meat were homogenized with 15 mL of whole muscle solubilization buffer (2%wt/vol SDS, 10mM sodium phosphate buffer; pH 7.0) using a polytron (POLYTRON Kinimatica CH-6010, Switzerland) at medium speed for 5 s bursts for 30 s. Homogenate was filtered through a double-layered cheese cloth and 1.4 mL of homogenate was placed into an Eppendorf tube (2mL safe-lock tube; 02236352, Eppendorf AG, Hamburg, Germany). Tubes were centrifuged at 1,500 x g for 15 min at 25°C to remove traces of insoluble components. 1 mL of supernatant was collected and into a 1.5 mL eppendorf tube. Protein concentration was determined using a Pierce bicinchoninic acid protein assay kit (Pierce Biotechnology, Rockford, IL, USA). All samples were diluted to 6.4 mg protein /mL using whole muscle solubilizing buffer. Samples were diluted to a final concentration of 4mg protein/mL with 50% of gel buffer (3mM EDTA, 3% wt/vol SDS, 30% vol/vol glycerol, 0.001% wt/vol pyronin Y, and 30mM Tris-HCl; pH 8.0) and 10% of betamercaptoethanol and then heated on a block for

15 min at 50°C. 30µL of samples were loaded on 4-20% Mini-PROTEAN TGX™ precast polyacrylamide gels (Bio-Rad Laboratories, Hercules, CA, USA) with a 10µL pre-stained standard (Precision Plus Protein Kaleidoscope, #1610375, Bio Rad, Hercules, CA) using a Bio-Rad Mini-PROTEIN 2 Cells system (Bio-Rad Laboratories). The system was run at constant voltage of 360V for 45 min with an electrophoresis buffer (1xTris/Glycine/SDS, #161-0732, Bio-Rad Laboratories, Hercules, CA, USA). Proteins in the gels were blotted to polyvinylidene difluoride membranes (0.45µm, Immobion-FL transfer membrane; Millipore) using a Bio-Rad Mini-Trans-Blot Electrophoretic transfer cell (Bio-Rad Laboratories) for 90 min at a constant voltage of 90V with ice-cold transfer buffer (25mM Tris-base, 192mM Glycine, 20% methanol; pH at 9.2). Membranes were blocked for 60 min in Odyssey Blocking Buffer (LI-COR, Lincoln, NE, USA) and incubated for 60 min at room temperature in monoclonal anti-desmin antibody (DE-U-10 Sigma-Aldrich, St. Louis, MO, USA) at a dilution of 1:10,000 in Odyssey blocking buffer containing 0.2% TWEEN-20 and 5% non-fat dry milk. Membranes were incubated overnight at 4°C, then washed three times with Tris Buffered Saline containing 0.2% TWEEN-20 (TBST) for 10 min and incubated in IRDye 680 LT Conjugated Goat Anti-Mouse IgG1 secondary antibody (LI-COR) at a dilution of 1: 10,000 in Odyssey blocking buffer containing 0.2% TWEEN-20 for 60 min. Membranes were washed three times with TBST and scanned using Odyssey Infrared Imaging system (LI-COR) at 700nm. Degradation was evaluated by quantifying band intensities (k. pixels) using Odyssey application software version 1.1. Bands at 55kDa were designated as intact and bands at 38kDa were designated as degraded. Percent degradation was calculated by (intensity of degraded bands/intensity of intact bands)* 100.

Free calcium concentration

Free calcium was quantified according to the procedure described by Parrish et al. (1981) with modifications. Three grams of powdered sample were centrifuged at 196,000 x g (Beckman Optima XPN-90 Ultracentrifuge, Type 50.2 Ti rotor, Beckman Coulter, Brea, CA) at 4°C for 30 min. Seven hundred µL of the supernatant were collected and treated with 0.1 mL of 27.5 trichloroacetic acid (TCA). Samples were centrifuged at 6,000 x g (accuSpin Micro 17R, ThermoFisher Scientific, Waltham, MA) for 10 min at 4°C. Four hundred µL of supernatant were transferred to a syringe, and the volume was brought to 4 mL with deionized, distilled water. The diluted sample was filtered through a 13 mm diameter Millex-LG 0.20 µm syringe filter (Millipore, Bedford, MA). Calcium concentration was quantified at Ward Laboratories (Kearney, NE) using an inductively-coupled plasma emission spectrometer (iCAP 6500 Radial; Thermo Electron, Cambridge, UK) with an appropriate calcium concentration standard.

Isoprostanes

All analysis were completed using OxiSelect™ 8-iso-Prostaglandin F2alpha ELISA Kit (Cell BioLabs, INC., STA-337, San Diego, CA). Thirty mg of powdered meat were weighed in a 5 mL eppendorf tube. Samples were homogenized in 2 mL of 2N NaOH using a Micropolytron (Pro-Scientific200, Bio-Gen, Oxford, CT) and spun for 20 s, cleaning micropolytron between each sample. Homogenized samples were heated in a water bath (Thelco 261, GCA Precision Scientific, Chicago, IL) at 45°C for 2 h to ensure hydrolysis. Samples were cooled to room temperature (~20 min), then pH was neutralized using 2 mL of 2N HCl, then vortexed for 20 s. Two 2 mL Eppendorf tubes

(2mL safe-lock tube; 02236352, Eppendorf AG, Hamburg, Germany) were used to split samples. Using microcentrifuge, samples were spun at 10,000 x g for 15 min at 4°C. One hundred μ L of sample were added to 100 μ L of neutralization solution in a 0.6 mL Eppendorf tube. Additionally, Anti-8-iso-PGF2 α Antibody (1:1000) was diluted with sample diluent. One hundred μ L of the diluted antibody to the Goat Anti-Rabbit Antibody coated plate and incubated on an oscillation shake at 25°C for 1 h. During incubation, 100 mL of 1x wash buffer was prepared by diluting 10x wash buffer concentrate with deionized water. Isoprostanes standards were prepared as labeled in Appendix XV. After incubation, antibody solution was removed from the wells and washed 5 times with 300 μ L 1x wash buffer per well. After the last wash, wells were emptied and tapped on absorbent pad or paper towel to remove excess wash solution. Immediately before use, conjugate (1:80) was diluted with sample diluent. Fifty-five μ L of the 8-iso-PGF2 α standard or hydrolyzed sample were combined with 55 μ L of 8-iso-PGF2 α -HRP conjugate in a microtube and mixed thoroughly. One hundred μ L of the combined solution per well were mixed thoroughly. Using a well containing sample diluent as a blank, microwell plates were incubated for 1 h at 25°C on an orbital shaker. After incubation, combined solutions were removed from the wells and washed 5 times with 300 μ L of 1x wash buffer per well. After the last wash, emptied wells were tapped on absorbent pad or paper. One hundred μ L of substrate solution was added to each well and incubated at 25°C for 30 min on an oscillate shaker. After incubation, 100 μ L of stop solution was added to each well. Results were read immediately on a microplate reader using 450nm as the target wavelength. Units of isoprostane content were designated as picogram/mL.

Sarcomere length

Sarcomere length was determined using the helium-neon laser diffraction method described by Cross et al. (1981) and Dolazza and Lorenzen (2014). A few flecks of powdered meat sample were placed on a clear glass microscopic slide. A single drop of 0.25M sucrose solution was added to the slide and topped with a glass coverslip. The distance to the top of the slide from the base of the laser was 100mm. A sheet of paper was placed below the stand in order to mark the two diffraction bands. Six sarcomeres per sample were determined and sarcomere length (μm) was determined by the equation provided by Cross et al. (1981):

Sarcomere length (μm) =

$$\mu = \frac{0.6328 \times D \sqrt{\left(\frac{T^2}{D}\right) + 1}}{T}$$

Where:

0.6328 = 632.8 (the wavelength of the laser) $\times 10^{-3}$

D = distance from specimen to diffraction pattern screen (100mm)

T = spacing between diffraction bands (mm)

Proximate composition

Moisture, fat, and ash (%) of raw meat samples were determined. Samples were measured in triplicate in Whatman #2 paper filter paper and fat was extracted with anhydrous ether. After identifying and recording weights of the folded filter papers with corresponding paper clips to hold packets closed, these were tared out and powdered meat samples (2 g) were weighed onto the filter paper and then closed with the paper

clip. Samples were then placed in a Soxhlet tubes and the boiling flasks were filled with 400 mL of ether. Once in place, water was opened to enter the condensers and each individual burner was turned on. After 48 h, burners were turned off and allowed to cool completely. Samples were air-dried under a fume hood for 2 h, then placed in a drying oven (105°C) overnight prior to recording final dry weight. In order to calculate final fat percentage, the final equation was used: % Fat = ((Pre-extraction wet weight with filter paper and paper clip – Post-extraction dry sample weight)/sample weight) * 100) - % Moisture. Moisture and ash (%) were measured with a LECO Thermogravimetric Analyzer in duplicate (Model 604-100-400, LECO Corporation, St. Joseph, MI), see Appendix IV for information. Moisture was determined in nitrogen atmosphere with a start temperature of 25°C and an end temperature of 130°C (17 min ramp rate). Ash was determined in oxygen atmosphere with a start temperature of 130°C and an end temperature of 600°C (30 min ramp rate). Protein was determined by difference.

Fatty acids

Fatty acid profiles were obtained via gas chromatography as described by (Folch et al., 1987). After extraction, lipids were converted to fatty acid methyl esters according to Morrison and Smith (1964) and Metcalfe et al. (1966). One g of powdered sample was homogenized with 5 mL of 2:1 chloroform: methanol and allowed to sit at room temperature (23°C) for 1 h. After, samples were filtered through Whatman #2 paper, brought up to a final volume of 10mL with 2:1 chloroform: methanol, and vortexed for 5 s with 2mL of 0.74% KCl. Samples were centrifuged (1,000 x g for 5 min at 5°C) and the top layer was aspirated off. After centrifugation, samples were dried on a heating block at

60°C under nitrogen purge. Once dry, 1 mL of 0.5 M NaOH in methanol was added, vortexed (5 s), and again heated at 100°C for 10 min. One mL of 14% Boron Trifluoride in methanol (wt/vol) was added, vortexed (5 s), and again heated at 100°C, this time for 5 min. Two mL of saturated salt solution and 2 mL of hexane was added and vortexed (5 s). Samples were centrifuged (1,000 x g for 5 min at 5°C) and the hexane layer removed and analyzed using gas chromatography (TRACE 1310 Gas Chromatograph; ThermoFisher Scientific, Waltham, MA). Fatty acids were separated using a Chrompack CP-Sil 88 capillary column (0.25mm by 100mm; Inlet temp: 260°C, Oven: 140°C hold for 5 min, increase at 4°C/min to 240°C and hold for 15 min. FID temp: 280°C. Injected at 30:1 ratio) and identified based on their retention times compared to known commercial standards (NU-Check Prep, Inc., Elysian, MN; #GLC-68D, GLC-79, GLC-87, GLC-455, and GLC-458). The percentage of fatty acids were determined by the peak areas in the chromatograph and values were converted to mg/100g tissue: Fatty acid mg/100g tissue = (% of fatty acid peak area * fat content of samples) * 100.

pH analysis

Powdered sample from chops from all aging periods with 0 d RD were weighed out in 5 g duplicates into 250 mL plastic beakers and placed on a stir plate. Forty-five mL of distilled deionized water and a magnetic stir bar were added to ensure constant mixing during the measurement process. The pH was measured using a pH meter (Orion 410Aplus; ThermoFisher Scientific; Waltham, MA) that was calibrated using 4.0, 7.0, 10.0 standards. The mean measurement of the duplicates was utilized for all analysis.

Objective color and subjective color in simulated retail display

Objective color measurements were taken once daily for 7 days during simulated retail display at all aging time points. Chops (2.54 cm) were placed on Styrofoam trays (21.6 x 15.9 x 2.1 cm, Styro-Tech, Denver, CO), overwrapped with oxygen permeable film (Prime Source PSM 18 #75003815, Bunzl Processors Division, North Kansas City, MO), and placed under retail display conditions (3°C under white fluorescence lightening at 1,000 to 1,800 lux). Commission international de l'éclairage (CIE) L* a* b* values were obtained using a Minolta CR-400 colorimeter (Minolta, Osaka, Japan) set with a D65 illuminant, 2°C, with an 8 mm diameter measurement area. Three measurements were made per chop and the mean was calculated for statistical analysis. The colorimeter was calibrated daily with a white ceramic tile (Calibration Plate, Serial No. 14933058, Konica Minolta, Japan). Lightness (L*) is measured with a range from 0 (black) to 100 (white), a* measures redness with the range between red (positive) and green (negative), and b* is a measure of yellowness from yellow (positive) to blue (negative). Color readings were recorded at the same time each day.

Visual discoloration was assessed daily during the 7 d of RD utilizing 5 trained panelists comprised of graduate students from the University of Nebraska. Panelists were trained using a standardized discoloration guide (Appendix VI). Discoloration % was approximated from 0% to 100% with 0% meaning no discoloration present and 100% being a fully discolored chop. Chops were randomly rotated daily to minimize location effects.

Lipid oxidation (TBARS)

Lipid oxidation was determined using thiobarbaturic acid reactive substances values (TBARS) for all aging periods at 0 and 7 d RD described (Ahn et al., 1998). Five grams of powdered meat were placed into a 50 mL conical tube to which 14 mL of distilled deionized water were added and 1 mL of butylated hydroxyanisole (BHA) solution (10% BHA: 90% ethanol). Samples were homogenized using a Polytron (POLYTRON® Kinimatica CH-6010, Switzerland) for 15 s at medium to high speed. The samples were centrifuged (2,000 x g for 5 min at 10°C) and one mL of supernatant was transferred into a 15 mL conical tube with 2 mL of 2,4,6-tribromoanisole (TBA) 2,4,6-tricholoroanisole (TCA) solution (15% and 20 mM TBA in deionized distilled water). Tubes were then placed in a 70°C water bath for 30 min. After 30 min, tubes were cooled for at least 10 min in a water bath (22°C) and centrifuged (2,000 x g for 15 min at 10°C). Two hundred µL of supernatant was transferred to a 96-well plate in duplicate (Microtest III sterile 96 well flat-bottomed microplate; Becton Dickinson & Company, Lincoln Park, NJ). Absorbance values were then read at 540 nm using a microplate spectrophotometer and compared to known standards (Model Epoch Biotek, Winooski, VT). Results were expressed in mg of malonaldehyde per kg of tissue.

Statistical analysis

Statistical analysis was conducted with SAS (version 9.4, Cary, NC). Objective and subjective color data were analyzed as a split-plot repeated measures design with treatment as the whole-plot, aging period as the split-plot and retail display as the repeated measures. Tenderness, troponin t, desmin, calcium, and pH were analyzed as a

split-plot design with treatment as the whole-plot and aging period as the split-plot. Lipid oxidation, free thiols and carbonyls were a split-split-plot design with treatment as the whole plot, aging period as the split-plot and retail display time as the split-split-plot. Transcriptomics, sarcomere length, fatty acids, cytochrome c, and isoprostanes were analyzed as a completely randomized design. Data were analyzed using the PROC GLIMMIX procedure of SAS and animal was the experimental unit. Correlations were evaluated using the PROC CORR procedure of SAS across all postmortem analyses. All means comparing within aging periods were separated using SLICE function in SAS. All means were separated using the LS MEANS statement with an α level of 0.05 and tendencies were considered at an α level of 0.10.

Results

Rectal temperatures

In this study, treatment had a significant effect ($P < 0.05$) on rectal temperatures of lambs (Figure 1), and a time effect ($P < 0.05$) was identified. Lambs administered LPS50 and LPS100 had significantly greater ($P < 0.05$) rectal temperatures than lambs administered the saline control. Additionally, LPS-treated lambs exhibiting the greatest increase in rectal temperature at 1, 2, and 4 hours post-injection, with a numerically greater rectal temperature in LPS100 treated lambs compared to LPS50 at each timepoint listed. These results are in agreement with Yates et al. (2011), who found LPS treatments to consistently increase rectal temperatures and peak at 4 hours post-injection, followed by a steady decline to basal temperature around 24 hours.

Transcriptomics

Considering all transcripts with differential expression ($P_{\text{raw}} < 0.05$), pathway analyses identified 68 conical pathways altered due to treatments. Transcriptomics pathway analyses are illustrated in Figure 2, with total change in gene turnover expressed as negative log (-log). Additionally, all significant ($P < 0.05$) pathways are expressed by a color-treatment directionality (Orange = LPS100; Blue = Control), with increased level of standard deviations (z-score) shown by an increase in the intensity of the respective colors. In the LPS100-treated samples, genes with evidence of dysregulation due to treatment were found in pathways that predicted an upregulation of genes responsible for cellular biosynthesis, oxidative stress generation, cellular defense systems, nucleic acid alteration, and skeletal muscle function.

Tenderness

Warner-Bratzler Shear Force (WBSF) for lambs across treatment and aging are shown in Figure 3. A significant aging effect ($P < 0.0001$) was identified across aging periods, along with a trend ($P = 0.10$) found across treatments. No aging time-by-treatment interaction ($P = 0.13$) was seen.

A significant aging effect ($P < 0.0001$) was found, as 1 day-aged chops exhibited greater WBSF compared to 14 day-aged chops (7.31kg and 2.52kg, respectively). Treatment tended to affect WBSF ($P = 0.10$), as chops from lambs administered LPS50 had lower shear force values compared to chops from lambs administered saline (Control) (4.51kg and 5.41kg, respectively), with no differences in shear force values between LPS100 and Control. No aging-time-by-treatment interaction ($P = 0.13$) was

seen. Interestingly, chops from lambs administered LPS50 had noticeably lower WBSF compared to Control (6.59kg and 8.06kg, respectively) within 1 d aging. There were no differences ($P > 0.05$) in tenderness between LPS100 and Control among aging periods, however, chops from lambs administered LPS100 had numerically lower shear force values than Control at 1 day of aging (7.27 kg and 8.06 kg, respectively). Chops aged for 14 days did not differ in WBSF across treatments ($P > 0.05$).

Troponin T

After 1 day postmortem, a treatment effect was found ($P = 0.02$), with LPS50 samples having significantly greater percent troponin-T degradation compared to Control and LPS100 (Figure 4) (10.32%, 6.85%, and 6.24%, respectively). No differences in degradation were seen at 14 days postmortem. A significant aging effect ($P < 0.0001$) was found, as 1 day-aged chops exhibited less troponin-T degradation compared to 14 day-aged chops. There was no aging time-by-treatment effect ($P > 0.05$). It is interesting, however, that chops from lambs treated with LPS50 had numerically greater degradation compared to Control and LPS100 at both 1-day aging and 14-day aging. These results parallel the WBSF data.

Desmin

In this study, a significant aging effect was found ($P < 0.0001$), as 1-day aged chops exhibited less percent desmin degradation compared to 14-day aged chops (3.57% and 50.6%, respectively; Figure 5). An aging-by treatment effect trend ($P = 0.08$) was found. Within 1-day aging, no differences ($P > 0.05$) were found across treatments.

However, both Control and LPS100 tended ($P = 0.10$) to be greater in percent degradation compared to LPS50 at 14-days aging (55.02%, 54.94%, and 41.73%, respectively).

Free calcium concentration

Free calcium concentration for samples across treatments and aging are shown in Table 1. An aging effect was shown for free calcium, as 14 day-aged samples had notably higher ($P < 0.0001$) free (sarcolemmic) calcium compared to 1 day-aged samples. No differences ($P > 0.05$) were found across treatments.

Isoprostanes

There were no differences ($P = 0.21$) across treatments for F₂-isoprostane content (Figure 6). Interestingly, LPS50 and LPS100 treated samples had numerically greater total F₂-isoprostane content compared to Control chops (239.51 pg/mL, 219.95 pg/mL, 165.51 pg/mL, respectively).

Sarcomere length

No differences in sarcomere length among treatments were observed ($P = 0.70$; Figure 7). Similar results were found by Starkey et al. (2015) who reported postmortem sarcomere lengths of 1.77 μm in lamb longissimus muscle.

Proximate analysis

In this study, treatment had no effect ($P \geq 0.31$) on proximate composition of 1 d aged chops (Table 2). The mean values for the proximate composition were: 75.35% moisture, 14.49% protein, 8.49% fat, 1.67% ash.

Fatty acids

There were no differences ($P \geq 0.21$) found in amount of saturated fatty acids, monounsaturated fatty acids, polyunsaturated fatty acids, or trans fats, saturated:unsaturated ratio, along with no differences ($P > 0.05$) in any individual fatty acids identified (Table 3).

pH

Data for pH across treatment and aging in lamb chops are exhibited in Figure 8. There was no treatment effect for pH ($P > 0.05$), regardless of aging period. An aging effect was found for pH ($P < 0.0001$), as 14 day-aged chops had a greater pH (5.87) compared to 1 day-aged chops (5.71). However, the average difference in values are likely of little practical significance. There was no treatment by aging interaction ($P = .7621$).

Color (objective)

Color values are shown in Table 4. A days of aging-by-retail display interaction was detected ($P < 0.05$) for all color measures. In general, L^* values increased and a^* and b^* values decreased as aging and retail display increased, regardless of treatment.

L* values were statistically different ($P = 0.0017$) among dietary treatments. Regardless of days of aging and retail display, L* values were greater (lighter color) in LPS100 compared to Control treatments (45.97 and 44.37, respectively). Chops from the LPS50 treatment was not different from the other treatments. There were no days of aging-by-treatment interaction for L* values ($P > 0.05$).

There was a days of aging-by treatment effect ($P = 0.0008$) in a* (redness) values. Regardless of retail display, chops from Control lambs exhibited the lowest a* value across treatments at 14 days of aging, with no difference when comparing LPS50 and LPS100 (15, 16.01, and 15.70, respectively). There were no significant differences found for a* at 1 day of aging. Concurrently, an age-by-treatment effect ($P = 0.02$) was observed in b* values. Within 1 day of aging, meat from lambs treated with LPS50 exhibited ($P < 0.05$) the highest b* values compared to all other treatments. There were no significant differences at 14 days of aging. However, chops from lambs treated with LPS100 exhibited the highest b* values compared to all other treatments.

Discoloration

As expected, percent discoloration of chops increased ($P < 0.0001$) as days of retail display increased (Table 5). A days of aging-by-treatment effect was shown in Table 4 ($P = 0.02$). Chops from Control lambs clearly had the most discoloration across all treatments at 14 days of aging ($P < 0.05$). There were no differences among treatments at 1 day of aging.

Thiobarbaturic acid reactive substances

No differences ($P > 0.05$) in treatment were identified (Table 6). As days of retail display increased, lipid oxidation (mg malonaldehyde) also increased ($P < 0.0001$). A days of aging tendency was found, as 14 day-aged chops tended ($P = 0.06$) to have more malonaldehyde content compared to 1 day-aged chops (2.26 mg and 1.65 mg malonaldehyde, respectively). While not significant ($P = 0.17$), chops aged for 14 days with 7 days of retail display (14-7) had the greatest TBARS values compared with all other age-by-retail display combinations (14-7: 4.08; 1-7: 3.03; 14-0: 0.44; 1-0: 0.27, mg malonaldehyde).

R² Correlations

Significant correlations ($P < 0.05$) were found in analyses comparing analyses relevant to tenderness, including WBSF and calcium, WBSF and pH, and calcium and pH (Table 6). Additionally, there were significant correlations among analyses related to oxidative potential, including TBARS and pH, TBARS and calcium, and calcium and pH. Interestingly, there were significant correlations found between isoprostanes content and early postmortem tenderness measurements, including WBSF and desmin. Additionally, statistical trends ($P < 0.10$) were found in comparisons relating tenderness attributes, including positive comparisons across WBSF, desmin, calcium, troponin T, and sarcomere length.

Discussion

LPS treatment and oxidative response mechanisms

This research explored the relationship between oxidative stress response via LPS exposure, and the subsequent impacts of oxidative processes on meat quality. There is a wide variety of commercially-available LPS preparations and the type of LPS used in an analysis dictates the degree of physiological stress induced. Therefore, effects due to LPS treatment and its impact on oxidative stress are rather complex (Suliman et al., 2004).

Lipopolysaccharides are covalently-bound lipids derived from the outermost membrane of many gram-negative bacterium (*Escherichia coli*). Acting as an endotoxin, LPS promotes an acute inflammatory response (Sternberg, 2007; Powers et al., 2011). As a result, innate immune response mechanisms are activated, such as nuclear factor kappa-light-chain-enhancer of activated B cells (NF κ B), start transcription of pro-inflammatory cytokines interleukin-1 β (IL-1), interleukin-6 (IL-6), and tumor necrosis factor- α (TNF- α) (Halawa et al., 2013). Similar to hormones like cortisol, cytokines are broad proteins which function for cellular signaling of homeostatic mechanisms during biological stress.

Given the complexity of oxidative stress *in vivo*, administration of lipopolysaccharides provided a dose dependent method to quantitatively induce oxidative stress to monitor response factors. The LPS model (O111:B4) promoted a quick inflammatory response *in vivo*, as indicated by the rapid change in rectal temperatures of lambs (Yates et al., 2011). Under conditions of high LPS concentration, production of pro-inflammatory mediators such as ROS are produced as a result of LPS toxicity, inducing oxidative stress and altering composition of muscle cells, including apoptotic mechanisms (Suliman et al., 2004). While recent literature (Niu et al., 2016) has used

LPS as a means to generate oxidative stress and influence meat quality, the impact of reintroducing LPS every 72 hours in an immune challenge, as our study performed, has yet to be determined. We postulated that the constant reintroduction of an oxidative promoter would provide a basal level of oxidative stress prior to harvest, replicating the effects of conditions linked as potential oxidative promoters in livestock production systems (e.g., genetics, diet, environment, handling, etc.) which contribute to final meat quality.

In addition, the development of ROS can impact gene expression in muscle cells, promoting changes in cell proliferation, differentiation, function, and turnover. Using proteomic analysis, Malheiros et al. (2019) observed greater damage from oxidative stress in tender beef, compared to tough beef. The proteomic profile identified oxidation to antioxidant enzymes, heat shock proteins, and structural proteins, suggesting that biochemical changes within muscle induced by oxidative stress prior to harvest can impact meat tenderness. In the present study, transcriptomics was utilized to evaluate changes in RNA transcripts among samples induced by a LPS challenge. Lambs administered LPS100 exhibited an upregulation of genes related to stress response pathways compared to the Control. This was evident by the gross changes in systematic pathways responsible for cell biosynthesis and turnover (ILK, Ceramide, IGF-1, PI3K, EGF, ERK5). These pathways are predominantly involved in cellular metabolism, proliferation, differentiation, protein transfer, and cell signaling. Interestingly, certain pathways act as hybrid models, both acting for cell biosynthesis but have secondary functions to inhibit apoptosis (PI3K). This is rational, as a biological system that has undergone biological stress would endure cellular damage, and thus need mechanisms to

repair, grow, and reinforce new cellular structures, organelles, and signaling mechanisms. Conversely, Control samples also upregulated certain biosynthesis pathways (cAMP, ErbB2-ErbB3) compared to LPS100 samples, as homeostatic maintenance of cells requires continuous cellular turnover. It is noteworthy that genes upregulated in LPS100-treated lambs included those responsible high levels of cellular growth and differentiation, suggesting they can account for increased cellular turnover as a result of acute biological disruption. Additionally, LPS-treated lambs expressed nucleic-related proteins (HMGB1, Telomerase, Unfolded Protein, EIF2, Neurotrophin/TRK, JAK/Stat, 3-phosphoinositide). These proteins are responsible for the modification and development of chromosomes, chromatin function, and regulation of protein transcription, some of which are also capable of inducing cytokine signaling (JAK/Stat, HMGB1). In particular, any nucleic damage as a result of oxidative stress merits the degradation (Unfolded protein) of damaged proteins and renewal of nucleic information. Most notably, LPS-treated samples identified oxidative stress-related genes, all of which act for cellular oxidative response mechanisms (*IL-6*, *IL-8*, *IL-3*, *NRF2*, Chemokine), stress signaling (p38 MAPK, CXCR, TNFR-2, and Sumoylation), or autophagic initiation. Although overall statistical significance was fairly liberal ($P < 0.05$), the increased upregulation of systematic processes in response to LPS treatment in this study validates that oxidative stress was generated within our test subjects, allowing stress response systems to be evaluated.

Oxidative stress changed mRNA related to muscular function and development. In particular, LPS-treated samples exhibited an upregulation of NO/ROS generation in macrophages, neuromuscular signaling between neurons and myofibrils (Agrin), and

alterations in messenger molecules used to selectively block epithelial calcium channels (D-myo-inositol tetrakisphosphate biosynthesis pathways). This implies the induction of oxidative stress not only generated ROS, but also upregulated different proteins responsible for inhibition of calcium regulation. From the perspective of muscle contraction and early postmortem tenderization, these factors hold tremendous implications on the development of muscle function by neuromuscular signaling *in vivo* in addition to enzymatic action in post-rigor muscle tissue. Interestingly, genes upregulated in Control samples presented the opposite impact on muscle tissue. Specifically, an increase in calcium signaling was found, suggesting that a homeostatic environment facilitates increased availability of calcium release in myocytes. Additionally, an increase in PPAR α /RXR α , responsible for glucose and lipid metabolism, were identified in Control lambs. Lastly, eNOS signaling was upregulated in Control samples, reflecting support of endothelial homeostasis within the lining of blood vessels, in addition to maintenance of blood flow. Perhaps eNOS acts on mitochondrial biogenesis and fission within homeostatic environments (Tengan et al., 2012). This is plausible, as an increase in oxidative stress would induce mitophagy rather than mitochondrial fission. In total, these measures help to validate the presence of oxidative stress within our experiment, as well as identifying subsequent mechanisms which act in response to oxidative stress.

When discussing oxidative products, evaluation of biomarkers (isoprostanes) have recently been used as a measure of downstream products due to oxidative stress. F₂-isoprostanes are the family of prostaglandin-like compounds formed by non-enzymatic, free-radical oxidation of arachidonic acid (20:4) by reactive oxygen species. Due to an

integrated mechanism required to produce isoprostanes, its generation is contingent on ROS-mediated oxidation (Lawson et al., 1999; Milne et al., 2011). Generated during lipid peroxidation, F₂-isoprostanes are produced as esterified fatty acid form in phospholipids membranes, and then released using phospholipase action (Montuschi et al., 2004). Compared to other oxidative products, F₂-isoprostanes are very stable compounds that are detectable in all normal biological fluids and tissues. It is common to sample muscle biopsies or anoxic muscle tissue for isoprostane content, as the lack of blood present prevents F₂-isoprostanes to be transferred and metabolized in blood. Given the relationship between isoprostane generation and oxidative stress activity, detection of this molecule is regarded as the standard for evaluating oxidative stress. Isoprostanes has been linked to oxidative stress in animals due to its evaluation across a multitude of treatments speculated to influence oxidative stress *in vivo*, such as exercise Karamouzis et al. (2004). Ponnampalam et al. 2017) theorized the change in isoprostanes as an *in vivo* indicator of meat quality deterioration postmortem (*in vitro*). Using blood samples, they detected an increase in plasma of 8-isoprostane PGF₂α levels, with the feedlot diet exhibiting the greatest concentration at 4 and 8 weeks of feeding trials. This is due to both the increase in energy content of diets, but also the amount of α-tocopherol content increased in roughage-based diets (ryegrass, lucerne). Concurrently, positive correlations between 8-isoprostane PGF₂α levels and lipid oxidation (malonaldehyde content) were found. Interestingly, positive correlations between 8-isoprostane PGF₂α levels and both arachidonic (20:4) and linolenic (18:2n-6) acid were discovered. This supports Milne et al. (2011) on how F₂-isoprostanes can potentially be generated from other PUFAs [eicosapentaenoic acid (20:5n-3), adrenic acid (22:4n-6), and docosahexaenoic acid

(22:6n-3)] due to the orientation of their double bonds, but the knowledge on these fatty acids being able to generate F₂-isoprostanes is not fully understood. As a result, the generation of F₂-isoprostanes is an excellent marker for *in vivo* oxidative damage as a result of oxidative stress. Given our results, we did not find statistical evidence between Control and LPS treated samples. However, it is noted that LPS50 and LPS100 treated samples have noticeably greater isoprostane content compared to the control (~ 44.7% and 32.9% greater content, respectively). In conjunction with changes in RNA transcripts for genetic pathways, this supports our hypothesis that the presence of increased *in vivo* oxidative stress occurred in LPS treated samples.

Oxidative stress and meat tenderness

Tenderness is recognized as one of the most important factors for consumer palatability and repurchasing of meat (Koochmaraie and Geesink, 2006; Kemp et al., 2010). Subtle variations (~0.5 kg) in tenderness within muscles have been shown to greatly influence consumer likeness to different retail products (Martinez et al., 2017). While marbling acts as an indicator of tenderness (Emerson et al., 2013), a complete understanding of the biological mechanism of this relationship is not known. Such knowledge might be used to induce and increase the rate or extent of tenderization in meat. Oxidative stress has been linked to factors influencing meat quality, as oxidative stress has been suggested to impact organelles linked to normal muscle function and promote cellular turnover via apoptotic mechanisms (Stamler and Meissner, 2001; Kagan et al., 2009; Bolisetty and Jaimes, 2013). It has been postulated oxidative stress alters proteins responsible for muscle structure, color stability, and organelle components

interconnected with proteolytic mechanisms. As a result, oxidative stress could influence the degree of postmortem tenderization, and greatly impacting both product quality and consumer palatability (Ott et al., 2007; Mitacek et al., 2019; Xing et al., 2019).

This study's statistical evidence is not sufficient to claim LPS decreased WBSF. However, a tendency ($P = 0.10$) has been shown that LPS impacts tenderness of *longissimus* muscle early postmortem, as indicated by numerical differences in shear force at 1 day of aging. This tendency in early postmortem tenderization is reflected in the troponin-T degradation. Lonergan et al. (2001) indicated that degradation of troponin-T early postmortem (~2 days) could be used as a consistent indicator of proteolysis and muscle tenderness throughout muscle aging time up to 14 days aging. The action of oxidative stress catalyzes apoptotic events, possibly upregulating enzymatic degradation of proteins such as troponin-T (Sierra and Olivan, 2013). The experiment found a statistical increase in troponin-T degradation at 1 day postmortem for LPS50 samples, but not for LPS100 samples. Our analysis suggests that early postmortem degradation of troponin-T can be impacted under certain conditions of increased oxidative stress, influencing shear force and overall tenderness of meat. While these data are promising when hypothesizing an oxidative stress – tenderness relationship, no differences were observed for additional components typically used to explain meat tenderness differences (desmin, free calcium, sarcomere length, pH, proximate analysis).

It appears a lower level of oxidative stress induced by LPS50 was associated with lower shear force values compared to Control and LPS100 treatments. The statistical differences in troponin-T degradation at 1 day aging would support this change in WBSF early postmortem. Using much higher doses of LPS (3-6mg/kg) in poultry, Niu et al.

(2016) found greater WBSF in 1 day-aged samples than controls. In addition, the significant correlations found oxidative stress biomarkers (isoprostanes) to be positively associated with early postmortem tenderness and proteolytic degradation. Given this information, an argument can be made that different levels of LPS-mediated inflammation can induce different degrees of metabolic response across livestock species and therefore alter the impact on meat tenderness.

The influence of oxidative stress on meat is likely dependent on the timeframe in which samples are exposed to an oxidative stress promoter. From the literature, tenderness has been examined from samples administered oxidative stress pre- and postmortem. Cook et al. (1998) and (Wang et al., 2018) found lower shear force values when injecting pre-rigor *longissimus* muscle with nitric oxide promoters and H₂O₂ solution, respectively. It can be postulated that the overwhelming increase in reactive species exhausted antioxidant systems, and initiated expression of pro-apoptotic mechanisms (Bax, Bcl-2) responsible for organelle instability and mitophagy. Concomitantly, the degradation of organelle stability promotes efflux of molecular ions bound within organelles (cytochrome c), which facilitate calcium-dependent enzymatic degradation of proteins. Cottrell et al. (2008) identified increased tenderness in *longissimus* muscle when lambs were induced with nitric oxide inhibitors pre-slaughter. However, the same parameters resulted in higher shear force (less tender) values in *semitendinosus* samples. Suster et al. (2005) reported increasing the time to inject nitric oxide inhibitors prior to slaughter caused lower shear forces in both *longissimus* and *semitendinosus* muscles. These values paralleled increased myofibrillar fragmentation, an indicator of tenderness/proteolysis. Both experiments examined specific reactive species

in response to meat quality. Warner et al. (2005) postulated reactive nitrogen species (RNS), when working in concert with reactive oxygen species (ROS), can generate highly radical compounds capable of disrupting proteins responsible for organelle function pertinent to postmortem tenderization.

From this analysis, a multitude of parameters were examined which could influence oxidative stress. Factors such as metabolizable energy in animal diets, fatty acid composition, and structure of muscle fibers have all been implicated to impact meat tenderness, with evidence suggesting that oxidative stress may accentuate changes in meat quality by altering these factors (Chauhan et al., 2016; Starkey et al., 2016; Ribeiro et al., 2019). These assays were conducted to ensure there were no confounding factors with oxidative stress when assessing the impact on tenderness.

The results from this study suggest that LPS-mediated oxidative stress has the potential to model increased early postmortem tenderness, which is likely due to the intricate relationship between the prolonged generation of oxidative stress and *in vivo* alterations amongst biological mechanisms responsible for muscle tenderization within the myofibril postmortem. We speculate a few possible parameters responsible for oxidative stress impact on meat tenderness, including the duration of oxidative stress *in vivo* and degree of oxidative stress induced by the concentration of oxidative stress promoter, as shown in this experiment. Additionally, the isolation and analysis of individual muscles, as well as any additional extrinsic conditions used in normal management strategies which could promote oxidative stress, such as diet, temperature, exercise, or genetic selection of livestock. All of these conditions have been used in the literature as conditions to evaluate the impact of oxidative stress on meat quality.

However, the degree in which they manufacture oxidative stress *in vivo* is likely variable. Given the complexity of oxidative stress, it was valuable for our experiment to utilize a system to induce a controlled amount of oxidative stress, while elimination of confounding variables which could impact meat quality.

Oxidative stress and stability of color and lipids (pH, isoprostanes)

Given the degree of biological stressors produced during oxidative stress, it is critical to evaluate conditions which are detrimental to meat quality (Xing et al., 2019). Muscle pH was not greatly influenced by LPS treatment, suggesting our treatments did not generate sufficient stress to exhaust glycogen supplies and increase ultimate pH in meat. Additionally, neither lipid nor protein oxidation were affected by LPS treatment. Correlation coefficients reinforced the relationship of lipid oxidation and pH, with some indication that lipid oxidation can coincide with increased proteolysis (troponin T, calcium). These data, along with color values, indicate that oxidative stress measures used were not detrimental towards color stability or lipid oxidation. With regards to our experiment, oxidative stress-induced samples tended to maintain color stability, exhibiting sustained redness (a^*) and increased lightness (L^*) in LPS treated chops compared to Control samples. This is supported by noticeably less discoloration in LPS-treated lamb chops after aging. Oxidative stress damage was not supported by lipid oxidation, as there were no noticeable changes in malonaldehyde content. Interestingly, a numerical increase in oxidative biomarker (8-isoprostanes) content was seen in the absence of lipid oxidative differences, suggesting that oxidative stress occurred in greater capacity in LPS-treated lambs, but not to a degree to promote detrimental effects on lipid

oxidation. This observation conflicts with (Mitacek et al., 2019), as prolonged aging promoted mitochondrial degeneration, depletion of color reducing enzymes, and decreased color stability in their study.

Conclusion

In conclusion, lipopolysaccharide-mediated oxidative stress triggered the onset of biochemical pathways responsible for muscle composition, proliferation, sustainability, and apoptotic mechanisms related to meat quality. Although there were few statistical differences in factors contributing to meat tenderization caused by LPS treatment, there was a trend that showed lambs subjected to lipopolysaccharides had lower shear force values early postmortem compared to the saline control group. These findings could be the result of greater proteolysis of troponin-T early postmortem in chops from lambs administered LPS50. While there were no detriments in lipid oxidation and color stability, the change in isoprostane content could indicate the increase in apoptotic mechanisms *in vivo*, increasing oxidative mechanisms taking place early postmortem, resulting in the onset of apoptotic and proteolytic events responsible for enhanced tenderness in aged meat.

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Tables

Table 1. Free calcium concentration (μM) of chops (*L. lumbrorum*) aged for 1 and 14 days from lambs administered saline control (0), 50, or 100ng/kg bodyweight of LPS. (n=29)

	$\mu\text{M Ca}^{2+}/\mu\text{g mitochondrial protein}$					
	Age			<i>P</i> -value		
	1	14	<i>SEM</i>	Trt	Age	Trt x Age
Control	46.72b	108.02a	4.2274	0.4955	<0.0001	0.88
LPS50	40.70b	104.51a	4.456			
LPS100	43.63b	103.12a	4.2272			

^{a-b}Within a row, means without a common superscript differ at $P < 0.0001$

Table 2. Proximate Composition of lamb chops aged for 1 day from lambs administered saline control (0), 50, or 100 ng/kg bodyweight of LPS. (n=29)

	Proximate Composition			
	Fat (%)	Moisture (%)	Ash (%)	Protein (%)
Control	8.18	75.1	1.59	15.13
LPS50	8.29	75.51	1.77	14.43
LPS100	8.99	75.45	1.66	13.91
<i>SEM</i>	0.98	0.198	0.095	0.989
<i>P</i> -value	0.8174	0.3134	0.4442	0.6828

Table 3. Amount¹ of Fatty Acids from lamb chops (*L. lumbrorum*) aged for 1 day from lambs administered saline control (0), 50, or 100ng/kg bodyweight of LPS. (n=29)

Fatty Acid	Dietary Treatments			P-value
	Control	LPS50	LPS100	
C 10:0	2.8	2.96	2.8	0.5146
C 12:0	8.11	4.71	9.08	0.549
C 13:0	1.41	0.96	0.002	0.5844
C 14:0	181.12	183.25	213.03	0.6693
C 14:1	14.54	13.17	16.72	0.7083
C 15:0	38.15	33.03	40.04	0.5734
C 15:1	72.22	64.09	63.96	0.8796
C 16:0	1754.75	1786.67	1980.12	0.7341
C 16: 1T	30.31	21.73	31.14	0.2356
C 16:1	152.23	149.3	182.38	0.442
C 17:0	156.03	134.15	151.33	0.6655
C 17:1	174.3	151.26	173.56	0.703
C 18:0	1099.64	1111.18	1117.99	0.9944
C 18:1T	308.13	293.64	352.97	0.6015
C 18:1	3195.42	3282.1	3543.1	0.7905
C 18:1V	117.31	125.53	139.87	0.5919
C 18:2T	26.38	30.49	34.56	0.3524
C 18:2	493.34	547.45	584.04	0.6917
C 18: 3w3	15.77	20.04	22.23	0.218
C 18: 3w6	5.82	6.72	6.99	0.7998
C 19:0	6.63	3.98	4.54	0.6645
C 20:0	0.87	0.52	5.55E-17	0.4054
C 20:1	35.2	36.45	37.92	0.9419
C 20:2	12.62	12.88	9.48	0.6035
C 20: 4w6	170.12	175.81	166.74	0.9781
C 20:5	0.53	1.2	2.78E-18	0.5108
C 22:0	10.92	11.34	9.97	0.8824
C 22:5	13.64	14.1	14.71	0.9647
C 22:6	0	1.09	0	0.3413
C 24:0	19.96	19.59	18.83	0.9686
C 24:1	0	1.64	3.33E-18	0.3413
Total	8126.97	8249.87	8940.64	0.8164
SFA	32889.07	3301.18	3560.29	0.8603

MUFA	4099.67	4138.91	4541.61	0.7737
PUFA	738.23	809.78	838.74	0.8157
Trans	364.82	345.86	418.67	0.5457

¹Amount (mg/100 g tissue) of fatty acid in powdered loin samples determined by gas chromatography

²C16:1T: Palmitoleic acid; C18:1T: Elaidic acid; C18:1V: Vaccenic acid; C18:2T: Linolelaidic acid; C 18:3w3: α -Linolenic acid; C 18:3w6: γ -Linolenic acid; C 20:4w6: Arachidonic acid ;SFA: Saturated Fatty Acids; MUFA: Monounsaturated Fatty Acids; PUFA: Polyunsaturated Fatty Acids

Table 4. Objective color (L*, a*, b*) and Discoloration (%) of chops (*L. lumberum*) aged for 1 and 14 days from lambs administered saline control (0), 50, or 100 ng/kg bodyweight of LPS. (n=29).

Measure	Age	Treatment			P-value		
		Control	LPS50	LPS100	Trt	Age	Trt x Age
L*	1	44.37	45.47	45.92	0.0017	0.68	0.92
	14	44.36	45.73	46.06			
<i>Mean</i>		44.37b	45.6ab	45.97a			
a*	1	13.70a	13.67a	13.31a	0.01	<.0001	0.0008
	14	15.00b	16.01a	15.7a			
b*	1	6.93c	8.20a	7.39bc	0.12	0.12	0.02
	14	7.88a	7.55a	8.04a			
Discoloration	1	7.81a	3.34a	9.27a	0.35	0.22	0.02
	14	16.43a	3.32b	5.58b			

^{a-c}Means within the same row for a single trait are different ($P < 0.05$)

^{a-b}Overall means within the same row for a single measure different superscripts are different ($P < 0.05$)

L*: SEM Control (0) = 0.2747; SEM LPS50 = 0.55; SEM LPS100 = 0.3256

a*: SEM Control (0) = 0.1611; SEM LPS50 = 0.1698; SEM LPS100 = 0.1611

b*: SEM Control (0) = 0.1788; SEM LPS50 = 0.3435; SEM LPS100 = 0.2066

Discoloration: SEM Control (0) = 1.2415; SEM LPS50 = 1.3086; SEM LPS100 = 1.2415

Table 5. Percent Discoloration (%) across retail display.

Day	0	1	2	3	4	5	6	7	P-value
Discoloration %	0.03e	0.13e	0.57de	1.67de	4.41d	9.95c	18.66b	27.64a	<0.0001

^{a-c}Means denote significance between day ($P < 0.05$), SEM Day = 1.4598

r ² Coefficients														
	WBSF1	WBSF14	Troponin1	Troponin14	Desmin1	Desmin14	Calcium1	Calcium14	8-Iso	SL	pHI	pHI4	TBARS1	TBARS14
WBSF1	1	0.12675736	0.08215102	0.00124962	0.10736763	0.01288225	0.01304415	0.07177041	0.142884	0.00055037	0.02873703	0.03470769	0.08711352	0.05121169
WBSF14		1	0.03319684	0.01339112	0.02857114	0.09337302	0.00985056	0.20595259	0.00301173	0.01996286	0.15161678	0.066481466	0.07826566	0.06857066
Troponin1			1	0.01107967	0.12302556	0.00609961	0.00175059	0.1072235	0.02343961	0.00163458	0.01949933	0.000861423	0.04216041	0.03087049
Troponin14				1	0.00332006	0.05218483	0.04044925	0.05276209	0.00053685	0.05317636	0.08116231	0.112929603	0.18594206	0.18471944
Desmin1					1	0.06717427	0.00413835	0.00297243	0.29447988	0.0401281	0.00013596	0.00002601	0.00544792	0.02136567
Desmin14						1	0.01787302	0.0563825	0.00076066	0.00622837	0.01853682	2.48004E-05	0.0135839	0.00091264
Calcium1							1	0.09145786	0.06212058	0.01973463	0.0002042	0.156950669	0.0003316	0.00150389
Calcium14								1	0.04559079	0.10953452	0.15238093	0.308635803	0.36945731	0.21599256
8-Iso									1	0.00416025	0.00297352	0.023119203	0.00014256	0.00859514
SL										1	0.04347225	0.004647149	0.01722656	0.03960896
pHI											1	0.442437826	0.15353075	0.168141
pHI4												1	0.31740829	0.4430633
TBARS1													1	0.78834865
TBARS14														1

r² coefficients p < 0.10 p < 0.05

Table 6: Correlation Coefficients across all postmortem analyses.
WBSF: Warner-Bratzler Shear Force; 8-Iso: F₂-8 Isoprostanes; SL: Sarcomere Length; TBARS: Thiobarbaturic Acid Reactive Substances

Figures

Legend

Figure 1. Rectal Temperatures of lambs administered intravenous injections of Control, LPS50, or LPS100. Superscripts denote statistical differences ($P < 0.05$) within day.

Figure 2. Transcriptomics expressed by Conical Pathways in Con vs 100ng LPS treated lambs. P-values for negative logarithmic (-log) expression set for ($P_{\text{raw}} < 0.05$).

Figure 3. Warner-Bratzler shear force (WBSF) of chops (L. lumborum) from lambs administered Control, LPS50, or LPS100. Superscripts within the same aging period are different ($P=0.10$ Day 1: *SEM* Control = 0.3694, *SEM* LPS50 = 0.3894, *SEM* LPS100 = 0.3694; $P = 0.90$ Day 14: *SEM* Control = 0.1264, *SEM* LPS50 = 0.1332, *SEM* LPS100 = 0.1264)

Figure 4. Troponin-T degradation of chops (L. lumborum) aged 1 and 14 days from lambs administered Control, LPS50, or LPS100. Superscripts within the same aging period are different ($P = 0.02$ Day 1: *SEM* Control = 1.02, *SEM* LPS50 = 1.08, *SEM* LPS100 = 1.02; $P = 0.78$ Day 14: *SEM* Control = 7.90, *SEM* LPS50 = 8.33, *SEM* LPS100 = 7.90)

Figure 5. Desmin degradation of chops (L. lumborum) aged 1 and 14 days from lambs administered Control, LPS50, or LPS100. Superscripts within the same aging period are different ($P = 0.85$ Day 1: *SEM* Control = 1.41, *SEM* LPS50 = 1.49, *SEM* LPS100 = 1.41; $P = 0.10$ Day 14: *SEM* Control = 4.66, *SEM* LPS50 = 4.92, *SEM* LPS100 = 4.66)

Figure 6. Total F₂-8 isoprostanes content of chops (L. lumborum) aged 1 day from lambs administered Control, LPS50, or LPS100. ($P = 0.2053$); *SEM* Control = 29.0413, *SEM* LPS50 = 30.6122, *SEM* LPS100 = 29.0413)

Figure 7. Sarcomere Length of lamb chops (L. lumborum) aged for 1 day from lambs administered Control, LPS50, LPS100. Superscripts within the same aging period are different ($P > 0.05$; *SEM* Control = 0.02378, *SEM* LPS50 = 0.02507, *SEM* LPS100 = 0.02378)

Figure 8. pH of lamb chops (L. lumborum) for 1- and 14-days aging from lambs administered Control, LPS50, LPS100. Superscripts within the same aging period are different ($P > 0.05$; Day 1: *SEM* Control = 0.03, *SEM* LPS50 = 0.03, *SEM* LPS100 = 0.03; Day 14: *SEM* Control = 0.02, *SEM* LPS50 = 0.025, *SEM* LPS100 = 0.02)

Figure 9. Malonaldehyde content of lamb chops (L. lumborum) for 1- and 14-days aging from lambs administered Control, LPS50, LPS100. Superscripts within the same day per aging period are different ($P > 0.05$; *SEM* Control = 0.5491, *SEM* LPS50 = 0.5788, *SEM* LPS100 = 0.5491)

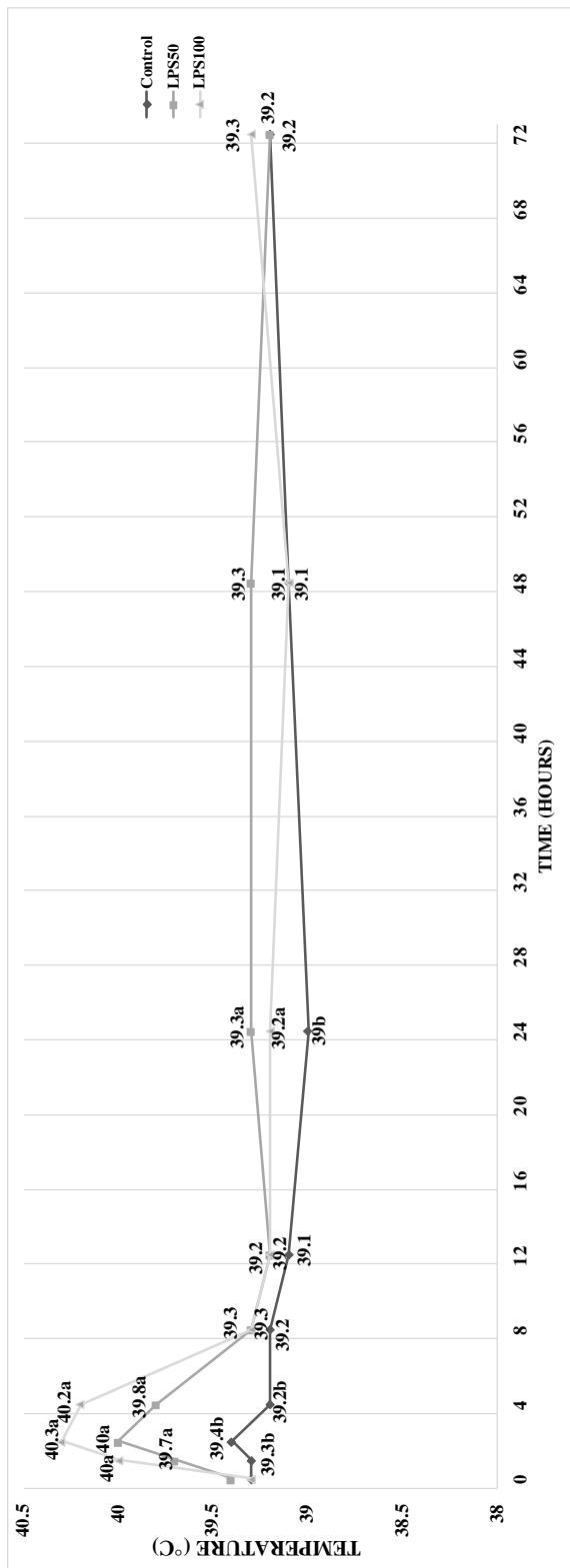


Figure 1. Rectal Temperatures of lambs administered intravenous injections of Control, LPS50, or LPS100.

^{a-b}Within a time denotes significant difference ($P < 0.05$).

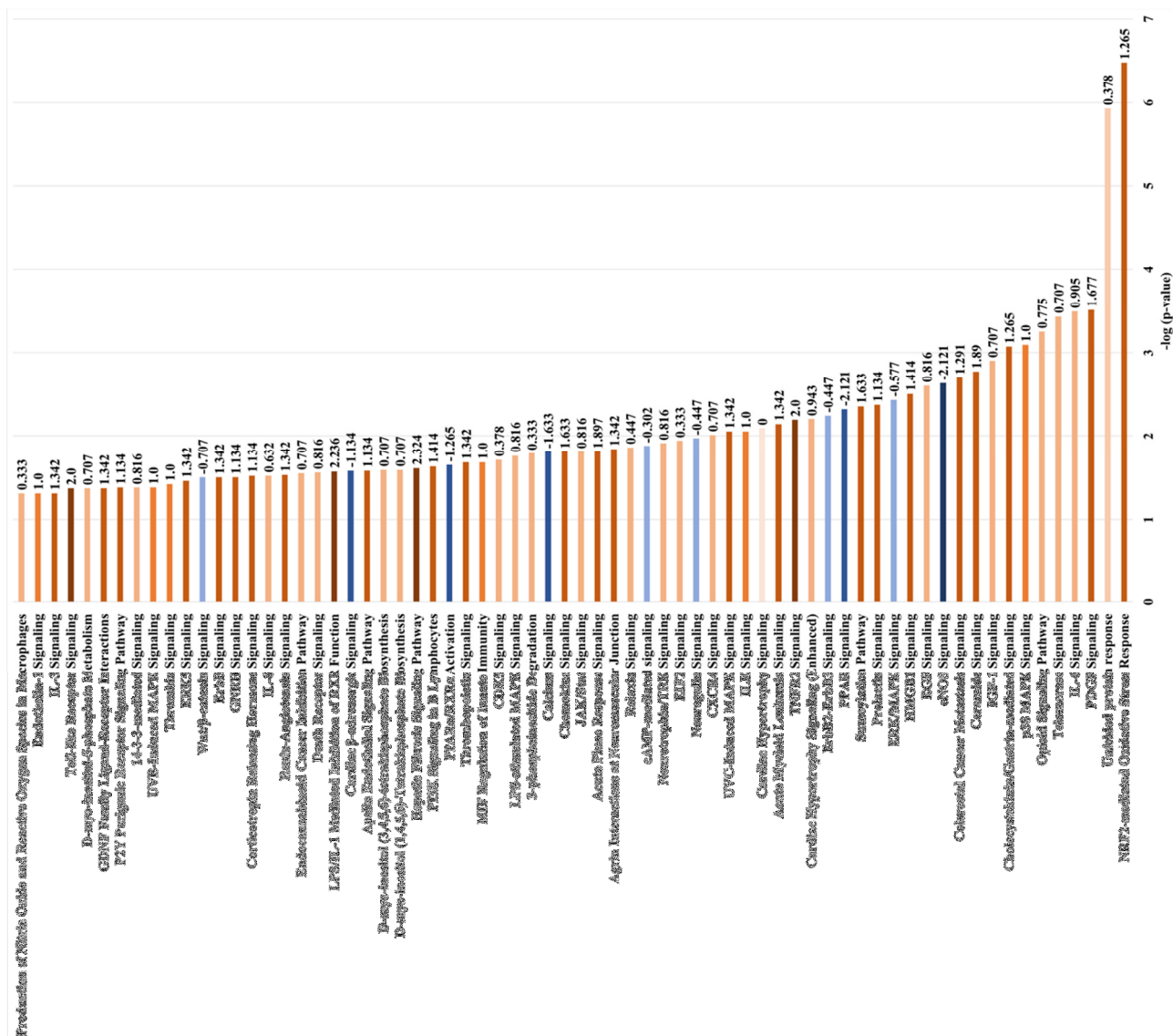


Figure 2: Conical Pathway Expression between Con and 100ng LPS treatments. Pathway Expressions are statistically significant ($P < 0.05$).
 Color Scheme: Orange – Upregulated to 100ng LPS; Blue – Upregulated to Con
 Numerical Values denote degree of z-score (Positive = Upregulated to 100ng LPS; Negative = Upregulated to Con)

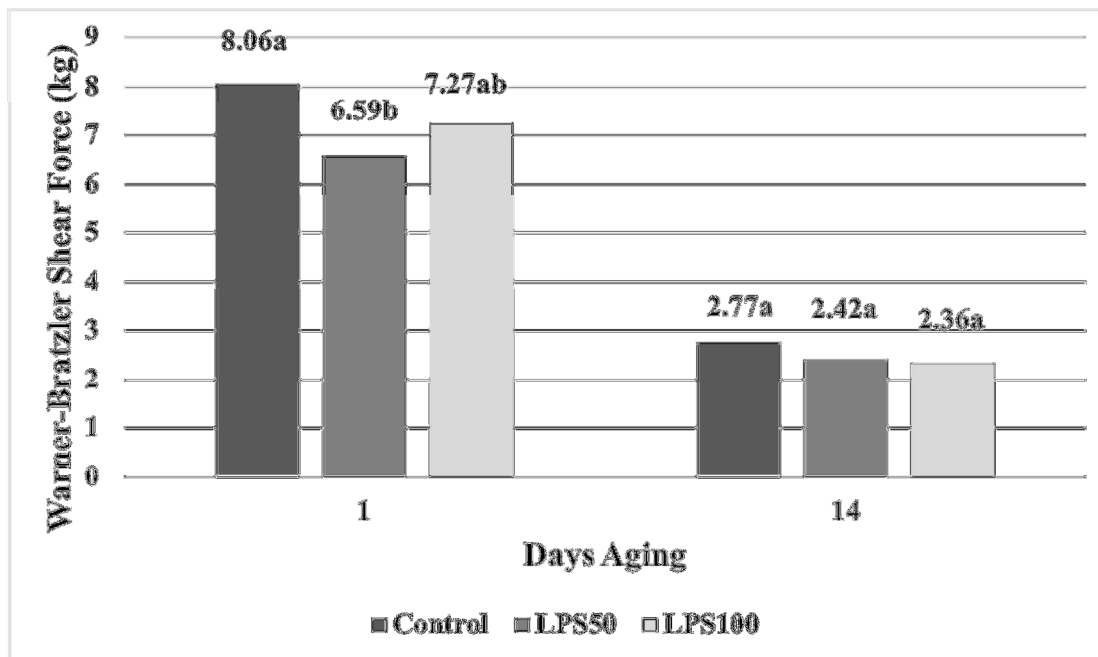


Figure 3. Warner-Bratzler shear force (WBSF) of chops (*L. lumbrorum*) from lambs administered Control, LPS50, or LPS100. Superscripts within the same aging period are different ($P=0.10$ Day 1: SEM Control = 0.3694, SEM LPS50 = 0.3894, SEM LPS100 = 0.3694; $P = 0.90$ Day 14: SEM Control = 0.1264, SEM LPS50 = 0.1332, SEM LPS100 = 0.1264)

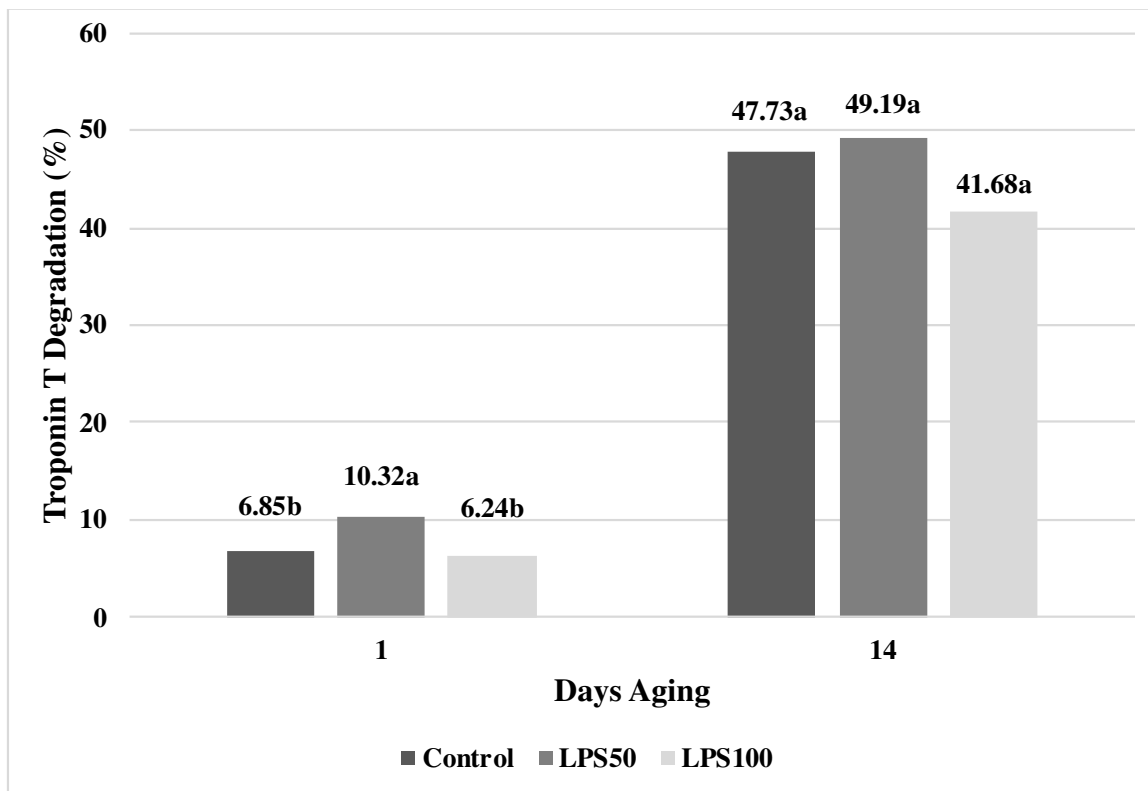


Figure 4. Troponin-T degradation of chops (*L. lumbrorum*) aged 1 and 14 days from lambs administered Control, LPS50, or LPS100. Superscripts within the same aging period are different ($P = 0.02$ Day 1: SEM Control = 1.02, SEM LPS50 = 1.08, SEM LPS100 = 1.02; $P = 0.78$ Day 14: SEM Control = 7.90, SEM LPS50 = 8.33, SEM LPS100 = 7.90)

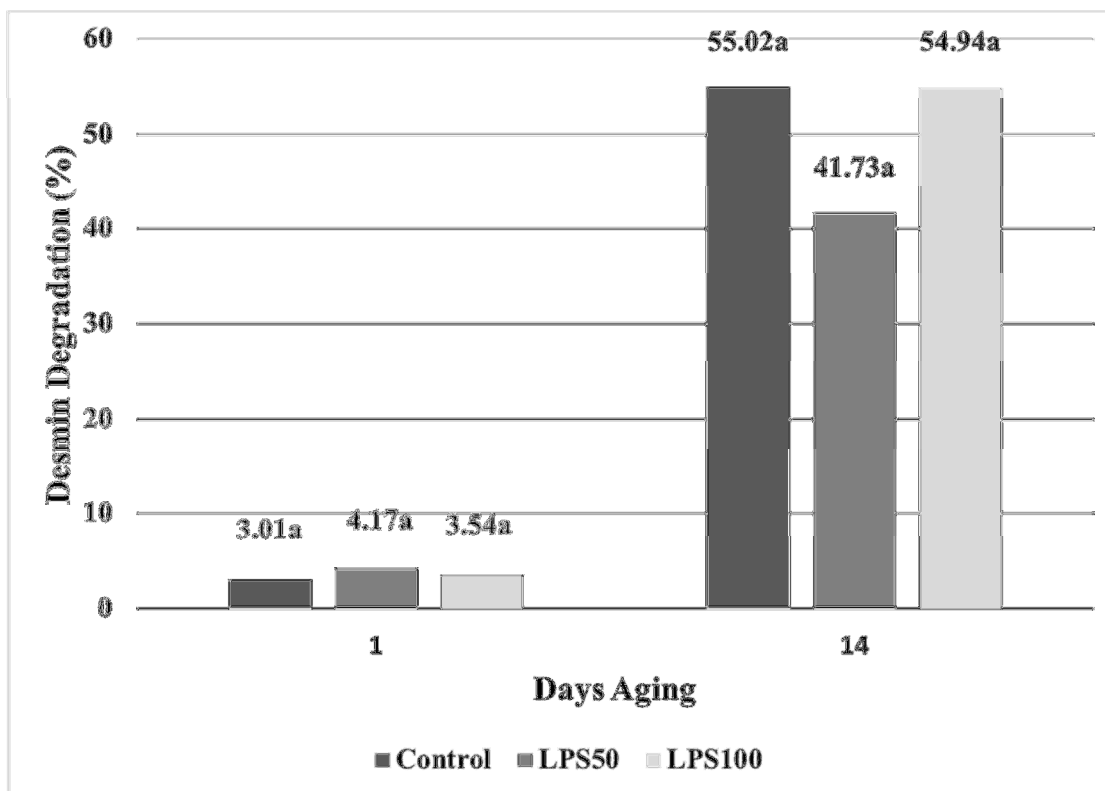


Figure 5. Desmin degradation of chops (*L. lumbrum*) aged 1 and 14 days from lambs administered Control, LPS50, or LPS100. Superscripts within the same aging period are different ($P = 0.85$ Day 1: SEM Control = 1.41, SEM LPS50 = 1.49, SEM LPS100 = 1.41; $P = 0.10$ Day 14: SEM Control = 4.66, SEM LPS50 = 4.92, SEM LPS100 = 4.66)

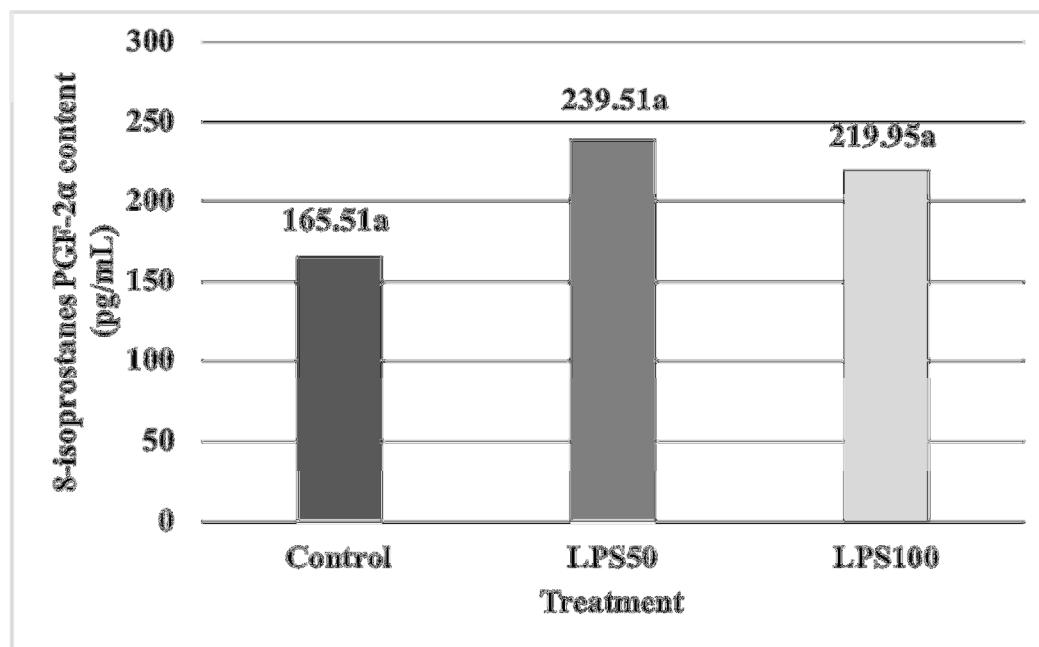


Figure 6. Total F₂-8 isoprostanes content of chops (*L. lumorum*) aged 1 day from lambs administered Control, LPS50, or LPS100. ($P = 0.2053$); *SEM* Control = 29.0413, *SEM* LPS50 = 30.6122, *SEM* LPS100 = 29.0413)

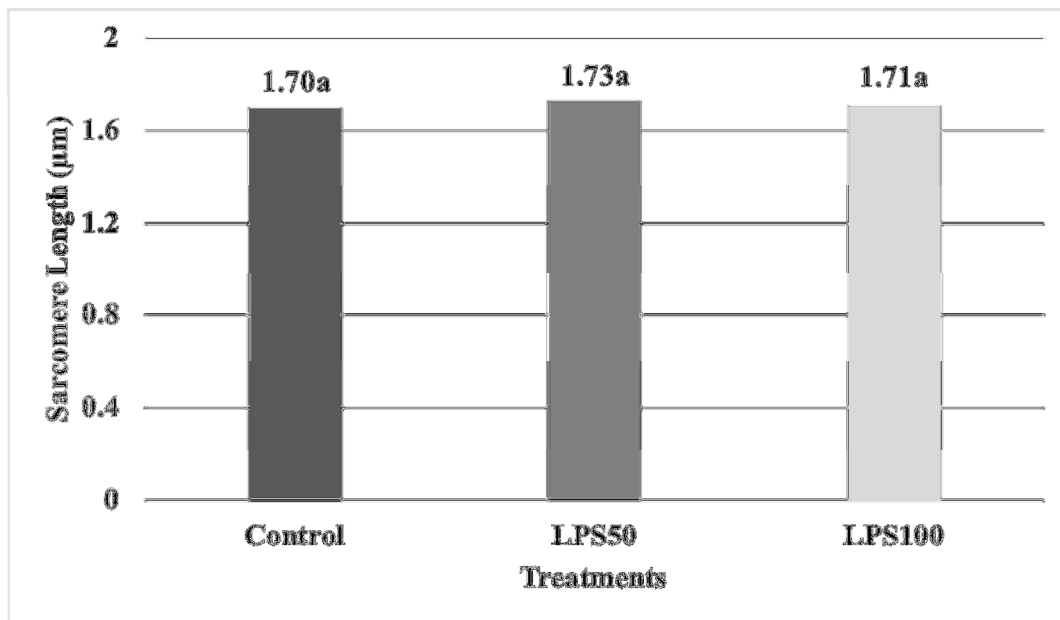


Figure 7. Sarcomere Length of lamb chops (*L. lumbrorum*) aged for 1 day from lambs administered Control, LPS50, LPS100. Superscripts within the same aging period are different ($P > 0.05$; SEM Control = 0.02378, SEM LPS50 = 0.02507, SEM LPS100 = 0.02378)

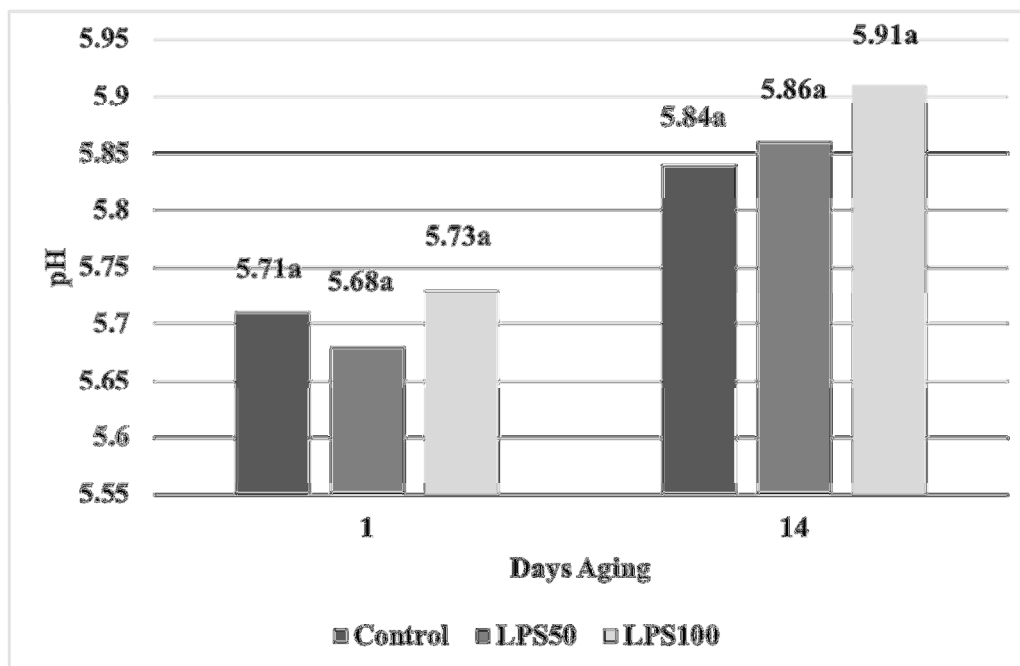


Figure 8. pH of lamb chops (*L. lumbarum*) for 1- and 14-days aging from lambs administered Control, LPS50, LPS100. Superscripts within the same aging period are different ($P > 0.05$; Day 1: *SEM* Control = 0.03, *SEM* LPS50 = 0.03, *SEM* LPS100 = 0.03; Day 14: *SEM* Control = 0.02, *SEM* LPS50 = 0.025, *SEM* LPS100 = 0.02)

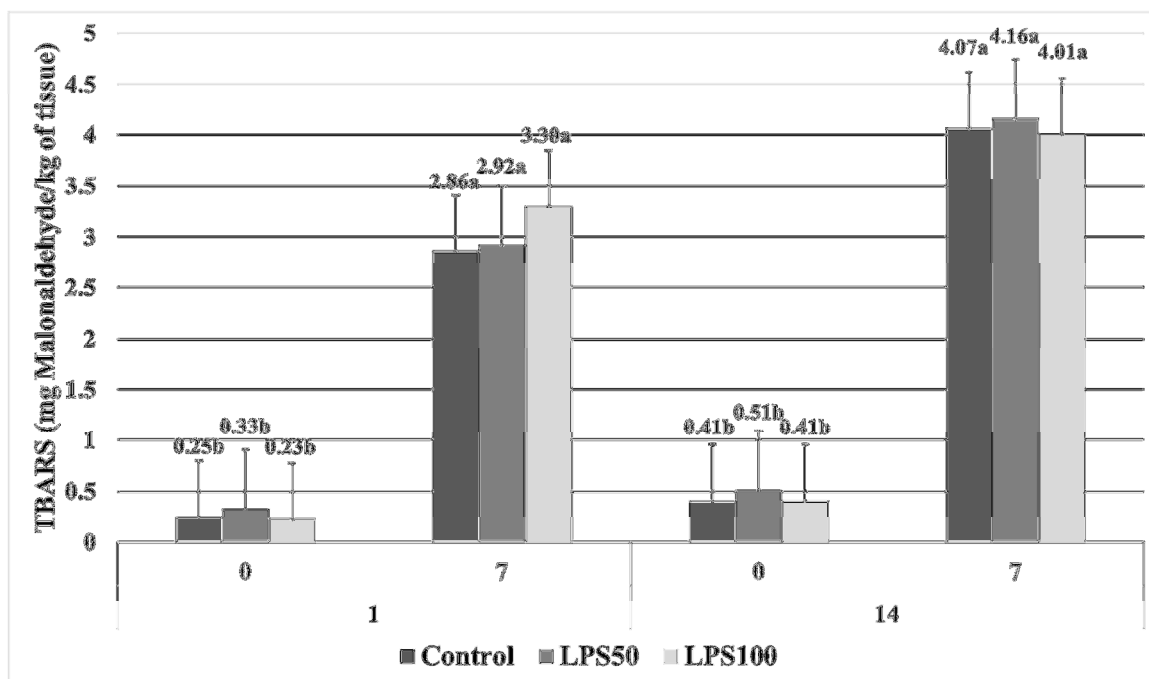


Figure 9. Malonaldehyde content of lamb chops (*L. lumbrorum*) for 1- and 14-days aging from lambs administered Control, LPS50, LPS100. Superscripts within the same day per aging period are different ($P > 0.05$; SEM Control = 0.5491, SEM LPS50 = 0.5788, SEM LPS100 = 0.5491)

RECOMMENDATIONS FOR FUTURE RESEARCH

Given the complexity of oxidative stress, there is a multitude of methods which could be used to evaluate its impact on meat quality. When analyzing *in vivo* oxidative stress on meat quality, I would recommend further exploration in the degree of oxidative stress by varying levels of lipopolysaccharide administration, as well as evaluate the amount of time in which animals are exposed to an oxidative stress challenge. Furthermore, I would explore the generation of oxidative stress markers in conjunction with known factors associated with oxidative stress in livestock, such as diet inclusions, housing techniques, and genetic backgrounding. Additionally, I would recommend the evaluation of oxidative stress by profiling the composition of reactive species generated within muscle tissue using a set of conditions known to induce oxidative stress. Finally, I would compare the overall impact of oxidative stress across a multi-muscle analysis. Given the vast differences in histochemical and morphological characteristics across muscles, an inherent change on the impact of meat quality due to oxidative stress is likely, and could expand our knowledge on the impact of oxidative stress factors and their relationship to pre- and postmortem mechanisms responsible for meat quality.

APPENDIX

Appendix I: Lamb Diet Composition*

Ingredient	Unit (lb/Ton)	Composition (%)
Cracked Corn	1051	52.55
Soybean Hulls	400	20
Oats	300	15
Soybean Meal – 47%	100	5
Liquid Molasses	100	5
Limestone	20	1
Salt	10	0.5
Urea	10	0.5
Ammonium Chloride	8	0.4
Vitamin A-D-E	0.2	0.01

Diet Nutritional Composition	
Crude Protein	13.57%
Non-Detergent Fiber	24.28%
Total Digestible Nutrients	8485%
Calcium	0.56%
Phosphorus	0.34%
Potassium	0.78%

*-Formulation Developed by Eastern Nebraska Research and Extension (1071 Co Rd G, Ithaca, NE 68033)

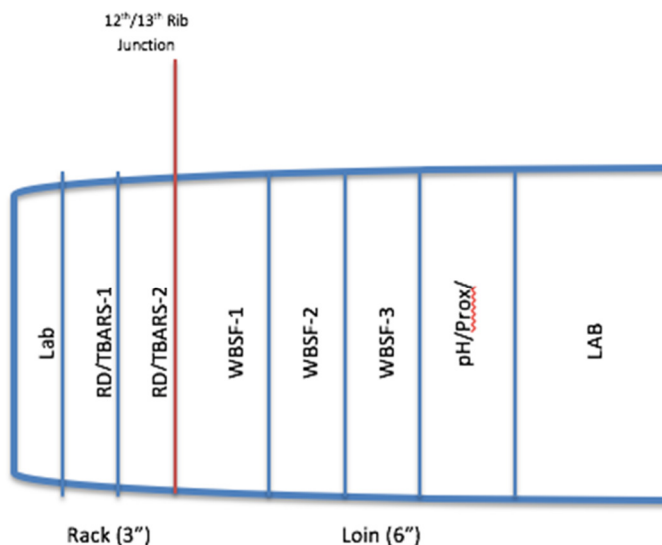
*-Low quality brome hay was given to lambs daily as a 1% of total body weight to ensure proper gut maintenance.

Appendix II: Fabrication Map

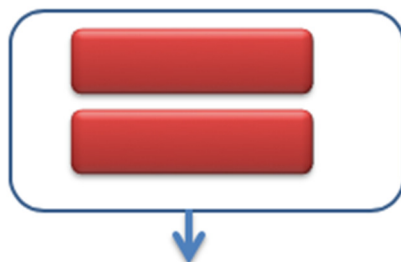
Fabrication map:

NOTES:

- Fabrication map for 1 and 14 d age
- WBSF and Lab chops (rib) will be 1"
- Extra Lab chops (loin) will be 2"
- Extra Lab chops (rack) will be 1"



Foam trays for RD:



Trays for **Color/ Discoloration:**

Each tray could have up to 2 one-inch chops, as long as they are of the same treatment (fat side inward)

20 chops per trt / 2 chops per tray = 10 trays

- 10 trays x 3 trt = 30 trays WBSF

Foam trays on RD per aging period = **30** trays per aging period

- Total foam trays = 30 x 2 aging periods = **60** Total Trays

NOTE: RD tables hold 35 trays (with proper lighting up to 40 trays). Therefore, we need 3 tables at every aging period.

Vacuum bags:

Size: 6 x 12

Lab samples: 32 animals x 2 aging periods x 2 lab samples = **128** bags

Size: 14 x 24

Loins: 32 samples x 1 (14 d age) = **32** bags**Additional supplies:**

- Powdering bags (~32 x 2(age) x 2(RD) = **128** bags)
- Liquid Nitrogen (~**4-5** tanks)

Tags needed:

- At the meat lab (Sample # - Age – Purpose – Replication):
 - UNL Carcass Tags = Set of 1-32
 - Ex.
 - UNL 1
 - **32** tags total
 - Cardstock for WBSF = Sets of 1-32 (laminated)
 - Ex:
 - 1-1-WBSF-1, 1-1-WBSF-2, 1-1-WBSF-3
 - 1-14-WBSF-1, 1-14-WBSF-2, 1-14-WBSF-3
 - **192** tags total
 - Cardstock for Aging = Set of 1-32 (**1 day aged samples will not require tags, as samples will be fabricated for WBSF and Lab samples**) (laminated)
 - Ex:
 - 1-14-Aging
 - **32** tags total
 - Cardstock for LAB samples = Sets of 1-32 (laminated)
 - Ex:
 - 1-1-LAB
 - 1-14-LAB
 - **64** tags total
 - Cardstock for LAB (Rib) Extra samples = Sets of 1-32 (laminated)
 - Ex:
 - 1-1-LAB-Xtra
 - 1-14-LAB-Xtra
 - **64** tags total
 - Color/Discoloration (adhesive tags) = Sets of 1-32
 - Ex:
 - 1-1-Color-7RD-1
 - 1-1-Color-7RD-2

- 1-14-Color-7RD-1
- 1-14-Color-7RD-2
- **128** tags total

Appendix III RNA Extraction

1. Heat Incubator to 55°C.
2. Clean work area with RNaseZap.
3. Add 10 μL of β -mercaptoethanol, per 1 mL of Buffer RLT before use.
 - a. Buffer RLT containing β -ME can be stored at room temperature (15-25 °C) for up to 1 month.
 - b. For 16 samples, 50 μL into 5 mL.
4. Clean weigh dish, tweezers and razor blade with 70% ethanol.
5. Pull tissue out of -80 °C freezer. Weigh $\leq 30\text{mg}$ tissue and put into 1.5mL tube.
6. Add 300 μL Buffer RLT solution. Vortex till broken up.
7. Homogenize with hand-held pestle for ~30 seconds.
8. Add 10 μL proteinase K solution.
9. Add 590 μL RNase-free water and mix thoroughly by pipetting.
10. Incubate at 55°C for 10 minutes.
11. Centrifuge for 3 minutes at 10,000 x g.
12. Transfer the supernatant (approximately 900 μL) into a new 2.0mL microcentrifuge tube (not supplied). Avoid transferring any of the pellet.
13. Add 0.5 volumes (usually 450 μL) of ethanol (96-100%) to the cleared lysate. Mix well by pipetting up and down. Do NOT centrifuge.
14. Transfer 700 μL of the sample to an RNeasy Mini spin column placed in a 2 mL collection tube. Centrifuge for 15 seconds.
15. Repeat step 7 using the remainder of the sample. Discard the flow-through.
16. Add 350 μL Buffer RW1 to the RNeasy spin column. Centrifuge for 15 seconds at $\geq 8,000$ x g to wash the membrane. Discard the flow-through.
17. Add 10 μL DNase I stock solution to 70 μL Buffer RDD. Mix by gently inverting the tube, and centrifuge briefly to collect residual liquid from the sides of the tube. *Note: DNase I is especially sensitive to physical denaturation. Mixing should only be carried out by gently inverting the tube. Do not vortex.*
18. Add the DNase I incubation mix (80 μL) directly to the RNeasy spin column membrane, and place on the benchtop for 15 minutes. *Note: Be sure to add DNase I incubation mix directly to RNeasy spin column membrane. DNase digestion will be in complete if part of the mix sticks to the walls or the O-ring of the spin column.*
19. Add 350 μL Buffer RW1 to the RNeasy spin column. Centrifuge for 15 seconds at $\geq 8,000$ x g. Discard the flow-through.
20. Add 500 μL Buffer RPE to the RNeasy spin column. Centrifuge for 15 seconds at $\geq 8,000$ x g. Discard the flow-through.

21. Add 500 μL Buffer RPE to the RNeasy spin column. Centrifuge for 2 minutes at $\geq 8,000 \times g$.
22. Place the RNeasy spin column in a new 2 mL collection tube (supplied), and discard the old collection tube with the flow-through. Centrifuge at 14,000 $\times g$ for 1 minute.
23. Place the RNeasy spin column in a new 1.5 mL collection tube (supplied). Add 30-50 μL RNase-free water directly to the RNeasy spin column membrane. To elute the RNA, centrifuge for 1 minute at $\geq 8,000 \times g$.

Appendix IV Objective color (L* a* b*) calibration instructions and helpful tips

Minolta Calibration Procedures

1. Before Calibration:

Calibrate Minolta in the same temperature conditions as the measurements being taken.

-Place the Minolta in the environment where samples will be measured about 5 to 10 minutes before calibrating so it can become equilibrated with the temperature.

Calibrate with the same materials as you will be taking measurements.

-If the measurement will not be taken directly on the meat surface, you must calibrate the Minolta with the same material it will be measuring through. For example, if you want to take readings from samples that are wrapped in overwrap, you must put some overwrap around the measuring head “eye” while calibrating using the white tile.

2. Turn the power to the measuring head *ON*.

3. Turn the power to the data processor *ON* while holding down the [DELETE/UNDO] key at the same time.

-Release the [DELETE/UNDO] key when you hear a BEEP. (This action deletes any previous data that might still be stored in the data processor)

4. When the screen turns on, the question “Initial set ok?” appears, press the [Measure Enter] key.

5. Once you get to the measurement screen, press the [Index Set] key.

a. Use the *arrows* and the [Measure Enter] key to adjust all the following settings:

- i. Printer: On
- ii. Color Space: Off
- iii. Protect: On
- iv. Auto Average: However many readings wanted per sample (1-30)
- v. Illuminant: D65
- vi. Back Light: Off
- vii. Buzzer: On
- viii. Disp. Limit

b. Press the [Esc] key to return to the measurement screen.

6. Press the [Calibrate] key while in the measurement screen.
7. Enter in the numbers listed on the calibrating white tile for the D65 setting using the following:
 - a. [←>] keys and the numeric pad
 - b. (The [←>] keys move the cursor)
 - c. D65 settings: Y: 93.13 x: 0.3164 y: 0.3330
8. Set up the measuring head so that it is resting on the LCD screen and the “eye” is facing up.
 - a. Place the white calibration tile on the measuring head, near the middle of the tile.
9. Press either the measurement button on the measuring head OR the [Measure Enter] key on the data processor after making sure the ready lamp is ON.
 - a. Make sure the white tile is completely on the measuring head “eye”.
 - b. The calibration is complete after the lamp flashes 3 times and the screen returns to the measurement screen.
 - c. Do not move the measuring head during calibration.
10. Press the [Color Space] key until the L* a* b* screen shows up.
11. Calibration is finished and the Minolta is ready.
 - a. To save battery life, turn both the measurer and data collector off after calibration is finished until you need it for measuring. The calibration and setting will not be erased.
 - b. When turning back on for measurements, ONLY turn on the power buttons. DO NOT hold down the [DELETE/UNDO] key at the same time. This will delete the calibration and settings and all of the steps will have to be repeated.

Cleaning:

- Wipe machine down with a soft, clean dry cloth. Never use a solvents such as thinner or benzene.
- If the white calibration tile becomes dirty, wipe it gently with a soft, clean dry cloth. If dirt is difficult to remove, wipe it with lens cleaner and cloth, then dry.

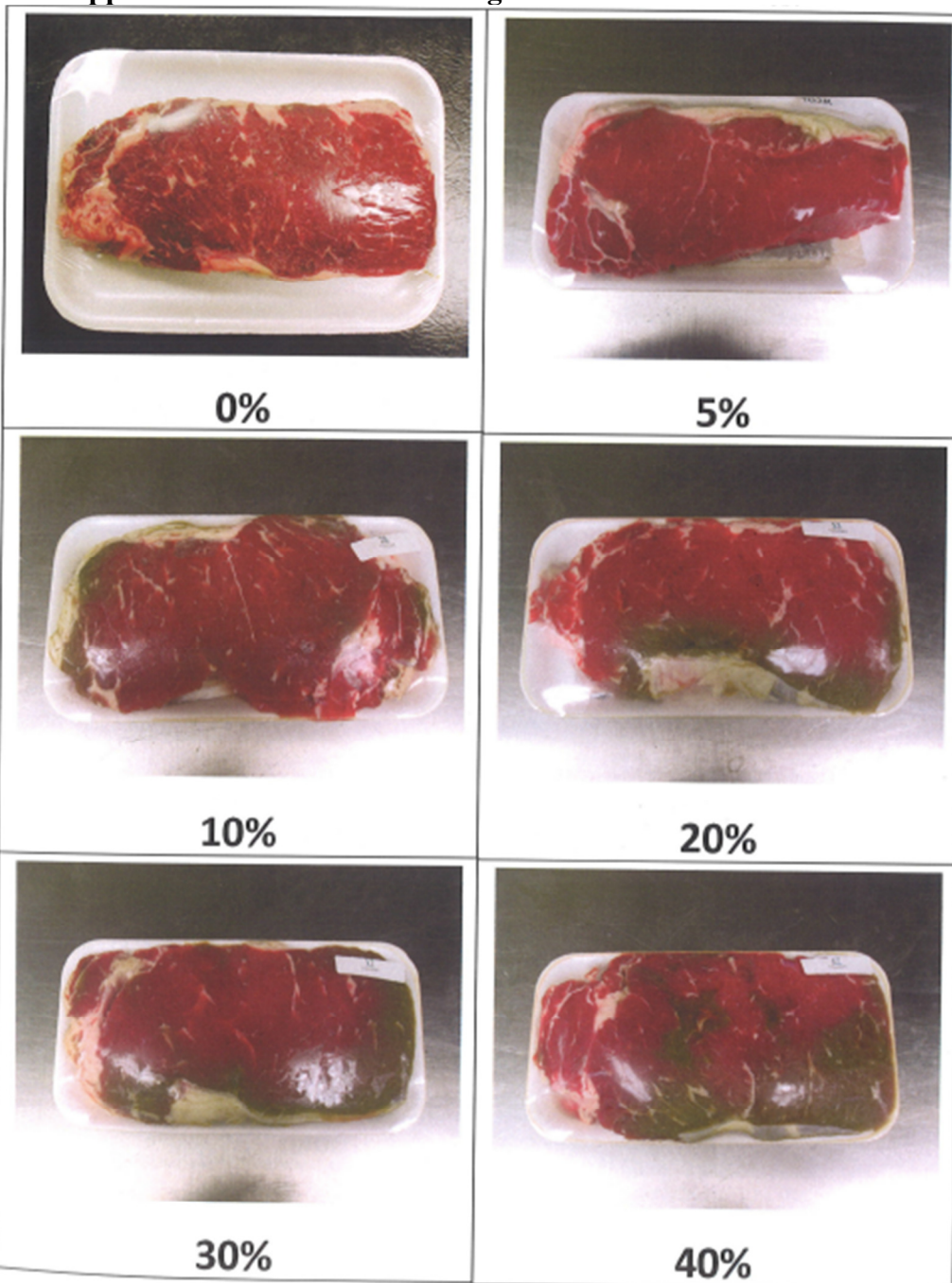
Minolta Helpful Tips

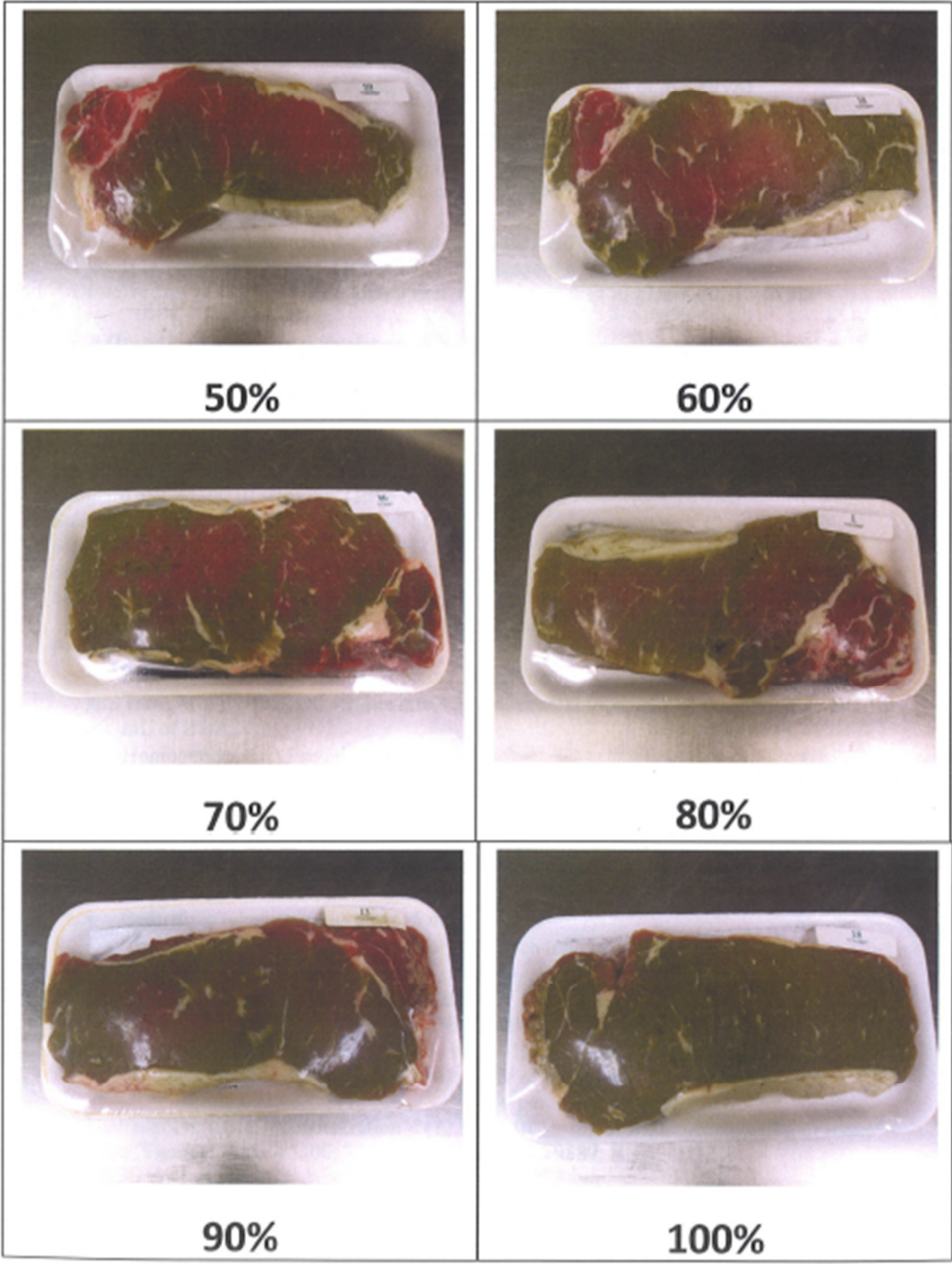
1. Make a separate data sheet
 - a. The Minolta prints out data with sample numbers 1 to 2000. In order to correlate it back to a sample, you must make a separate data sheet that has a place to record meat sample i.d. and its corresponding Minolta number.
2. Batteries

- a. The measuring head requires 4 AAA batteries and the data processor requires 4 AA batteries.
3. The auto protect setting
 - a. The Minolta can only record and store up to 2000 readings, once you go past 2000 readings it will start deleting older readings.
 - b. When the auto protect is on it will automatically prevent the 2001st reading from being taken so you cannot accidentally overwrite other data.
4. Auto Average Function
 - a. During calibration, if you set the Auto Average function to a reading number above 1, for example 5, you only have to hit the measure button once and it will automatically take all 5 readings then print out the average.
 - b. It only allows a second or two between readings so make sure you are paying attention and move the measuring head to where you want it before it automatically takes the next reading.
5. Recalibrate regularly
 - a. If using the Minolta all day, or for long periods of time, make sure to recalibrate it regularly.
6. DELETE/UNDO KEY
 - a. If you accidentally take a reading, hitting the [DELETE/UNDO] key will delete the last reading.
 - b. If you accidentally delete a reading by hitting the [DELETE/UNDO] key, hitting the [DELETE/UNDO] key again will restore the previous reading.
7. Printing Paper
 - a. The paper that the data is printed on is sensitive to heat and light. The printed data should be kept in a dark cool place, like a desk drawer. In order to prevent losing any data, you must make a photocopy of the printout in order to preserve it for long-term storage.
8. More than One Color Space on Print Out
 - a. If you want to print more than 1 color space (Example: L* a* b* and XYZ) on the print out slip:
 - i. Press the [Index Set] key. Use the arrows and the [Measure Enter] key to adjust all the following settings:
 1. Color Space: On
 2. Disp. Limit: press the [Measure Enter] key to select this option.
 3. Once inside the Disp. Limit option, go through the list and change all the color spaces that you DO NOT want to OFF.
 4. Press the [Esc] key until you return to the measurement screen.

9. Change Measurements to a Different Color Space
 - a. If you get done measuring and realize that you meant to measure in a different color space (For example: measured everything using Yxy and meant to use L*a*b*), you can correct it using these steps:
 - i. While in the measurement screen, press the [Color Space] key until your desired color space (in this example: L*a*b*) appears.
 - ii. Press the [Data List] key while in the measurement screen.
 - iii. Select the desired page using the up and down arrows. If you only have one page it will show up as P00, select this one.
 - iv. Once you have the desired page selected, press the [Measure Enter] key.
 - v. Press the [Print/Feed] key.
 - vi. Select “All Meas. Data” using the up and down arrows.
 - vii. Press the [Measure Enter] key. This will reprint all the stored data in your newly selected color space.
 - viii. Press the [Esc] key to return to the measurement screen.

Appendix V: Guide for Percentage of Surface Discoloration





Appendix VI

Fat extraction with Soxhlet method

Warning: Ether is extremely flammable and produces explosive peroxides. Never bring a radio or any other potentially spark-producing item into the fat extraction room.

1. Check ground glass connections. They should be wiped clean with a dry paper towel and given a thin coating of stopcock grease.
2. Each boiling flask must contain boiling stones. This helps prevent violent boiling of the solvent which could be dangerous.
3. Load samples into Soxhlet tubes, arrange them so that no samples are above the level of the top bend in the narrower tubing on the outside of the Soxhlet. (The Soxhlet will only fill with the solvent up to this point before cycling back down into the boiling flask.) In general, the large soxhlets will hold about 20 two-gram samples and the small soxhlets from 4-6.
4. Fill the large (500mL) boiling flasks with 400mL of solvent and the small (125mL) flasks with 100mL of solvent. Do this under the fume hood.
5. Fit the Soxhlet onto the boiling flask. Very carefully, bring the assembly into the extraction room and fit it onto the condenser. Make sure all ground glass connections are snug and each boiling flask is resting on the heating element. The ceramic fiber sheet should be covering the bare metal surfaces on the burners completely.
6. Turn on the water supply to the condensers (usually a quarter turn). Check later to make sure condensers are cool enough – if not, increase water flow.
7. Turn heating element control dials between three and four. Each burner has its own dial. Never turn the burner beyond five. Ether has a very low boiling point and violent boiling is dangerous. Double check fittings, boiling stones, etc.
8. Fat extraction will take from 24 to 72 hours depending on the sample. (Beef – 48 hours, Bacon – 72 hours). Check extractions twice daily to see that everything is alright while they are running.
9. When done, turn off the burners and let solvent cool completely before removing samples.
10. After it has cooled down, slowly uncouple the flask and Soxhlet tube from the condenser. Cover the top of the Soxhlet with one palm so as to reduce ether vapors while transporting it to the fume hood. Air dry samples in the fume hood for two hours to get rid of the remaining ether in the samples. Pour ether back slowly into an approved container for reuse or discarding. Do not leave ether out of the hood or the flammable cabinet.
11. Place samples in drying oven (105°C) for about four hours or overnight before weighing back.

Calculation: $\left(\frac{\text{Original weight including filter paper and paper clip} - \text{Fat extracted sample weight}}{\text{Sample weight}} \right) * 100 - \% \text{ Moisture} = \% \text{ Fat}$

Appendix VII

Minerals and Ash Determination

1. Place crucible in drying oven at 100°C for 4 hours and then in a desiccator.
2. Place 2 grams of pulverized muscle tissue into the crucible in duplicates.
3. Moisture and ash are determined with the following program:

Name	Covers	Ramp Rate	Ramp Time (minutes)	Start Temp (°C)	End Temp (°C)
Moisture	Off	6 d/m	17	25	130
Ash	Off	20 d/m	30	130	600

Name	Atmosp.	Flow Rate	Hold Time (minutes)	Constant Weight	Constant Weight Time (minutes)
Moisture	N	High	0	0.05%	9
Ash	O	High	0	0.05%	9

Crucible density and cover density were set at 3.00, while sample density was set at 1.00. The calculations to determine moisture and ash are the following:

$$\text{Moisture} = [(\text{Initial weight} - \text{Dry weight}) / \text{Initial weight}] * 100$$

$$\text{Ash} = (\text{Weight of ash} / \text{Initial weight}) * 100$$

Appendix VIII
Sarcomere Length of Powdered Meat Samples
(Dolazza and Lorenzen 2014; Cross et al. 1981)

1. Spread the powdered meat sample very lightly (just a few specks) on the microscope slide.
2. Place a drop of 0.25M sucrose solution on the slide and cover with a coverslip.
3. Place the slide on the stage of the laser stand (the distance between the slide and the base of the laser stand should be set to 100 mm).
4. Place a piece of paper at the base of the laser stand.
5. Move the slide back and forth through the laser light until a diffraction pattern is observed (Figure 1).

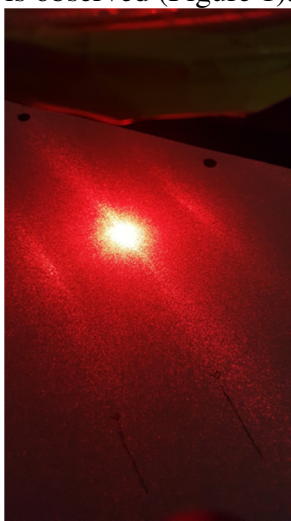


Figure 1. A projected sarcomere diffraction pattern on the paper.

6. Mark the original and the two diffraction bands for 5 different sarcomeres from the sample.
7. Measure the distances between the two diffraction bands and calculate sarcomere length using the following equation:

$$\text{Sarcomere Length } (\mu\text{m}) = \frac{0.6328 \times D \sqrt{\left(\frac{T}{D}\right)^2 + 1}}{T}$$

D = distance in mm from slide to the base of the laser stand.

T = ½ of the distance in mm from one first order band to the other first order band.

8. Use the average of the 5 sarcomere lengths to determine the sarcomere length of the samples.

Appendix IX

Fatty Acid Determination

1. Weight out 1 gram of pulverized muscle tissue. If extracting subcutaneous fat, weight out 0.1 gram of pulverized subcutaneous fat into centrifuge tube.
2. Add 5 mL of 2:1 chloroform: methanol (v/v) for muscle tissue for 3 mL for subcutaneous fat.
3. Vortex for 5 seconds and let stand for 1 hour at room temperature.
4. Filter homogenate through Whatman #2 filter paper into 13 x 150 mm screw cap tube bringing the final volume with chloroform: methanol to 10 mL for muscle lipid and 5 mL for subcutaneous fat extract. If stopping at this point, purge test tube with nitrogen, cap tube, and store at -80°C.
5. Add 2 mL of a 0.74% KCl solution for muscle lipid extract or 1 mL for subcutaneous fat tissue extract and vortex for 5 seconds. If stopping at this point, purge test tube with nitrogen, cap tube, and store at 0 °C for no more than 24 hours.
6. Centrifuge samples at 1,000 x g for 5 minutes. Following centrifugation, aspirate off the aqueous phase (top layer). If stopping at this point, purge test tube with nitrogen, cap tube, and store at -80°C.
7. Evaporate to dryness under nitrogen at 60°C.
8. Add 1 mL of a 0.5 M NaOH in methanol. Vortex for 5 seconds. Heat for 10 minutes at 100°C.
9. Add 1 mL of boron trifluoride in 14% methanol. Vortex for 5 seconds. Heat for 10 minutes at 100°C.
10. Add 2 mL of a saturated salt solution and 2 mL of hexane. Vortex for 5 seconds.
11. Centrifuge samples at 1,000 x g for 5 minutes. Following centrifugation, remove hexane layer (top layer) **making sure to not disrupt the aqueous phase** (lower layer) and place in GC vial. Purge GC vial with nitrogen, cap and crimp cap, and store at -80°C until sample is ready to be read on the GC.

GC Settings

Column- Chromopack CP-Sil 88 (0.25mm x 100mm)

Injector Temp- 270 °C

Detector Temp- 300 °C

Heat Pressure- 40 psi

Flow Rate- 1.0 mL/minute

Temperature Program- Start at 140 °C and hold for 10 minutes. Following 10 minutes, raise temperature 2 °C/minute until temperature reaches 220 °C. At 220 °C, hold for 20 minute.

Appendix X
Determination of Free-Calcium Level

(Parrish et al. 1981. J. Food Sci. 46:38-311 with modifications)

1. Analyze each sample in duplicates.
2. Mince frozen steak by using a stainless-steel knife (avoid fat and connective tissues).
3. Measure 3 g of minced meat into a thickwall polyallomer ultracentrifuge tube (13 x 55mm) and centrifuge at 196,000 x g for 30 min at 4 °C.
4. Pipette 700 μ L of the supernatant into an eppendorf tube.
5. Treat the supernatant with 0.1 mL of 27.5% trichloroacetic acid and vortex for 15 s.
6. After standing for 10 min at room temperature, centrifuge eppendorf tubes at 6,000 rpm for 10 min.
7. Pipette 500 μ L of the supernatant into a plastic tube and bring up the volume to 5 mL with deionized ddH₂O.
8. Filter prepared samples through a 13 mm diameter Millex-LG 0.20 μ m syringe filter (Millipore, Bedford MA) into a new tube.
9. Send prepared samples to the Ward Laboratories (Kearney, NE) for calcium level determination.
10. Calcium concentrations (ppm) of samples are quantified at the Ward Laboratories (Ward Laboratories, Kearney, NE) using the inductively-coupled plasma emission spectrometer (iCAP 6500 Radical; Thermo Electron Corporation, Cambridge, UK) with appropriate calcium concentration standards.
11. Calcium concentrations are calculated as follows:

Average ppm of Calcium in the sample	= X ppm = X mg/L
Molecular weight of Calcium	= 40.078 g
Micromolar (μ M) concentration of calcium	= (X x 1000)/40.078 μ M

Appendix XI Thiobarbituric Acid Reactive Substances Assay

TEP solution (1,1,3,3-Tetraethoxypropane)(Make new weekly)

Stock Solution: Dilute 99 μ L TEP (97%) bring volume to 100 mL ddH₂O.

Working Solution: Dilute stock solution to 1:3 (TEP Solution:ddH₂O)(1 x 10⁻³M).

TBA/TCA (2-Thiobarbaturic Acid/Trichloroacetic Acid) Stock Solution: 1L

15% TCA (w/v) and 20mM TBA (MW: 144.5) reagent in ddH₂O.

Dissolve 2.88g TBA w/warm ddH₂O first, then add TCA (150g) and ddH₂O to 1L.

BHA (Butylated Hydroxylanisole) stock Solution:

Make 10% stock solution by dissolving in 90% ethanol.

10 g BHA dissolved in 90 mL ethanol (90%) + 5 mL ddH₂O.

Standards: In duplicate

	<u>Moles of TEP</u>
Blank: 1 mL ddH ₂ O	
Standard 5: 100 μ L working TEP + 1.90 mL ddH ₂ O	(5 x 10 ⁻⁵ M)
Standard 4: 1 mL Std. 1 + 1 mL ddH ₂ O	(2.5 x 10 ⁻⁵ M)
Standard 3: 1 mL Std. 2 + 1 mL ddH ₂ O	(1.25 x 10 ⁻⁵ M)
Standard 2: 1 mL Std. 3 + 1 mL ddH ₂ O	(0.625 x 10 ⁻⁵ M)
Standard 1: 1 mL Std. 4 + 1 mL ddH ₂ O	(0.3125 x 10 ⁻⁵ M)

Remove 1 mL of Standard 1 and discard it, leaving 1 mL behind.

Procedure:

- Mix all reagent and standards before beginning.
- Transfer 5 g or powdered sample into a 50 mL conical tube; add 14 mL of ddH₂O and 1.0 mL of BHA.
- Homogenize for 15 sec with a polytron.
- Centrifuge for 2,000 x g for 5 min.
- Transfer 1 mL of homogenate or standard to 15 mL conical tube.
- Add 2 mL of TBA/TCA solution, vortex.
- Incubate in a 70 °C water bath for 30 min to develop color.
- Cool samples in a cold water bath for 10 min.
- Centrifuge tubes at 2,000 x g for 15 min.
- Transfer duplicate aliquots of 200 μ L from each tube into wells on a 96-well plate.
- Read absorbance at 540 nm.

Calculations: mg of malonaldehyde/kg of tissue

$$K(\text{extraction}) = (S/A) \times MW \times (10^6/E) \times 100$$

Where, S = Standard concentration (1 x 10⁻⁸) moles 1,1,3,3-Tetraethoxypropane)/5 mL

A = Absorbance of standard MW = MW of malonaldehyde (72.063 g/mole)

E = Sample equivalent (l) P = Percentage recovery

Final calculation: 0.012 x concentration x (72.063 x 10⁶) = mg of Malonaldehyde/kg of tissue

Reagents (Sigma): TBA- T5500; TCA -T9159; TEP – T9889; BHA – B1253

Appendix: XII
Isolation of Myofibrillar Proteins

(Pietrzak et al., 1997, J. Anim. Sci. 75:2106-2116)

1. Knife mince frozen steaks after trimming visible fats and connective tissues.
2. Weigh 3 g of minced meat into a 50 mL plastic conical tube.
3. Add 15 mL ice-cold rigor buffer (0.1M KCl, 2 mM MgCl₂, 1mM EGTA, and 10mM K₂HPO₄; pH 7.4) and homogenize using the polytron (POLYTRON Kinimatica CH-6010, Switzerland) at very low speed for 15 min.
4. Filter the homogenate through double-layered cheese cloth to remove connective tissue and fats into a new 50 mL plastic conical tube.
5. Pipette 1.4 mL homogenate into an eppendorf tube (2 mL safe-lock tubes; 02236352, Eppendorf AG, Hamburg, Germany).
6. Centrifuge eppendorf tubes at 4,000 x g for 5 min.
7. Decant the supernatant and dismantle the pellet using a cleaned spatula after re-suspending in 1 mL of ice-cold rigor buffer.
8. Vortex the mixture for 10 s and centrifuge for 5 min at 4,000 x g.
9. Repeat step 6 and 7 three times to remove myoglobin as much as possible (until the supernatant is clear and free of myoglobin).
10. Decant the supernatant and remove the leftover-supernatant using a pasture pipette.
11. Re-suspend the pellet in 250 μ L of ice-cold rigor buffer.
12. Vortex thoroughly after diminishing the pellet.
13. Store eppendorfs on ice for immediate use or in -80 °C for later use.

Reagent preparation:

- a. *Rigor Buffer (0.1 M KCl, 2mM MgCl₂, 1mM EGTA, and 10mM K₂HPO₄; pH 7.4)*

Add 14.91 g of potassium chloride (KCl: MW 74.55), 380.84 mg of magnesium chloride (MgCl₂: MW 95.21), 760.7 mg of ethylene glycol tetraacetic acid (EGTA: MW 380.35, and 3.484 g of potassium phosphate dibasic (K₂HPO₄): MW 174.18) into 1900 mL of deionized ddH₂O and dissolved properly. Check the pH and volume up to 2000 mL. (if want to adjust pH to 7.4, adjust using conc. HCl).

Appendix XIII

Troponin T Degradation

Protein Isolation

1. Weigh out 3 g powdered meat into a 50 mL plastic conical tube
2. Add 15 mL **rigor buffer**: 0.1 M KCl, 2mM MgCl₂, 1 mM EDTA, and 10 mM K₂HPO₄; pH 7.4.
3. Homogenize using polytron (POLYTRON Kinimatica CH-6010, Switzerland) at medium speed (setting 6) for 5 sec bursts until fully mixed
4. Filter homogenate through double layered cheese cloth into a new 50 mL plastic conical tube
5. Pipette 1.4 mL homogenate into an eppendorf tube (2 mL safe-lock tubes; 02236352, Eppendorf AG, Hamburg, Germany)
6. Centrifuge Eppendorf tubes at **4000 x g** for 5 min
7. Decant supernatant
8. Resuspend pellet in 1 mL rigor buffer
9. Vortex for 10 sec or until pellet is sufficiently mixed into solution
10. Centrifuge for 4000 x g for 5 min
11. Repeat 6-10 two more times
12. Decant supernatant fully
13. Suspend pellet in 1 mL **suspension buffer**: 0.1M Tris, 1.25mM EDTA, 5% SDS; pH 8
14. Vortex for 30 sec or until pellet is sufficiently mixed into solution
15. Centrifuge for 5 min at 4000 x g
16. Remove supernatant to new Eppendorf tube without removing remaining pellet.
This is the myofibrillar stock sample.

Protein Concentration

17. Dissolve 100 μ L of myofibrillar protein stock samples in 0.9 mL of suspension buffer (See step 13 in **Protein Isolation**) and vortex 10 sec
18. Prepare a concentration series (20-2000 μ g/mL) of bovine serum albumin (BSA) using suspension buffer
19. Place 25 μ L BSA standards and diluted myofibrillar protein samples on a microwell plate
20. Add 200 μ L BCA working reagents (50:1; Reagent A:Reagent B) into respective wells in the microwell plate
21. Mix protein samples and BCA working reagents thoroughly on a plate shaker for 30 sec
22. Incubate the microwell plate at 37°C for 30-40 min and cool to room temperature
23. Read absorbance at **562 nm** on a microplate reader (SpectraMAX 250, Molecular devices, Sunnyvale, California)
24. Protein concentrations are expressed as μ g/mL
25. Dilute protein to 2 μ g/ μ L with ddd water in new tubes

26. Dilute protein to 1 $\mu\text{g}/\mu\text{L}$ with Laemmli / betamercaptoethanol mixture: 950 μL Laemmli plus 50 μL betamercaptoethanol added.
27. Heat the tube on a heating block at 95°C for 5 min.

Gel Electrophoresis

28. Remove the Mini-PROTEAN TGX Precast Gel 4-20% from the package
29. Pull the tape off the bottom, remove the comb
30. Place into electrode assembly with short plate inward
31. Place second gel on the other end of the assembly
32. Lock the two gels in place
33. Place the assembly in the tank
34. Fill the chamber with Bio Rad 1x Tris/Glycine/SDS # 161-0732 running buffer until the short plate is completely covered
35. Add buffer to the outer tank to either 2 or 4 gel line, depending on how many gels you are running
36. Load 10 μL Bio-Rad Kaleidoscope Pre-stained standards in the first well
37. Load 5 μL sample into remaining wells
38. Place lid on tank, aligning color coded banana plugs and jacks
39. Set constant voltage to **200 V**
40. Run until tracking dye in the Laemmli buffer of each sample reaches the black finishing line (30 minutes to an hour)
41. Turn off power supply
42. Remove gel from assembly
43. Remove gel by gently opening key provided in precast gel package

Western Blot

44. Equilibrate gel in transfer buffer (25 mM Tris-base, 192 mM glycine, 20% methanol at pH 9.2) for 20 min
45. Soak precut filter paper (170-3932, Bio Rad Laboratories, Hercules CA) and fiber pads in transfer buffer for 5 min
46. Wet PVDF membranes (IPFL20200, Millipore) in methanol for 15 second and rinse briefly in deionized water before soaking in transfer buffer
47. Prepare the gel sandwich as follows:
 - a. Place the cassette with the black side (anode) down in the transfer buffer in the cassette assembly box
 - b. Place one pre-wetted fiber pad on the black side of the cassette
 - c. Place filter paper on the fiber pad
 - d. Place equilibrated gel on filter paper (roll out all bubbles)
 - e. Place pre-wetted membrane on the gel (roll out all bubbles)
 - f. Place the other filter paper (roll out all the bubbles) and fiber pad respectively
48. Close the cassette firmly without moving the gel and filter paper sandwich and lock the cassette with the white latch

49. Place prepared sets in module
50. Place module in tank and fill with transfer buffer
- 51. Run Western blot for 60 min at a constant amperage of 180mA**
52. To prevent nonspecific antibody binding, block membranes with 10 mL Odyssey Blocking buffer (927-40100; LI-COR, Lincoln NE) for 120 mins at room temperature

Primary Antibody Binding

53. Anti-Troponin-T (JLT-12: Sigma Aldrich, St. Louis, MO)
 - a. 1 μ L in 10 mL Odyssey Blocking Buffer + 0.2% Tween20 (20 μ L)
54. Incubate blots in diluted primary antibody for 60 min at room temperature while rocking
55. Incubate blots **overnight at 4°C** while rocking
56. Pour off primary antibody solution
57. Wash membrane four times for 10 min each with 15 mL of 1x Tris buffered saline + 0.1% Tween20 on rocking platform

Labeling Primary Antibodies

58. Reconstitute contents in the original IRDye 680LT conjugated goat anti-mouse IgG1 vial with 0.5 mL sterile distilled water
59. Mix gently by inverting and allow rehydrating for at least 30 min before use (this is stable for three months at 4°C) **DO NOT EXPOSE TO LIGHT**
 - a. 1 μ L in 10 mL Odyssey Blocking Buffer + 0.2% Tween20 (20 μ L)
60. Incubate blots in diluted secondary antibody for 60 min at room temperature with gentle shaking (make sure not to expose to light)
61. Pour off secondary antibody solution
62. Wash membrane four times for 10 min each with 15 mL of 1x Tris buffered saline + 0.1% Tween20 on rocking platform
63. Wash membrane with 15 mL of 1x Tris buffered saline for 30 mins to remove residual Tween20

Membrane Imaging

64. Use Odyssey Infrared Imaging System (LI-COR Biosciences, Lincoln NE) at 700 nm channel as integrated intensity
65. Imaging conditions:
 - a. Resolution: 169 nm
 - b. Quality: low
 - c. Focus: offset 0 mm
 - d. Channels: 700 nm
 - e. Intensity: 3.0 or 3.5
 - f. Image size: x-10 y-7

Appendix XIV

DESMIN

Whole-Muscle Protein Preparation

1. Weigh out of 400mg (0.4g) of powdered meat sample into a 50 mL plastic conical tube
2. Add 10mL of **Whole-muscle Solubilizing Buffer** (2%wt/vol SDS, 10mM sodium phosphate buffer, pH 7.0).
3. Homogenize using polytron (POLYTRON Kinimatica CH-6010, Switzerland) at medium speed (setting 6) for 5 sec bursts until fully mixed.
4. Filter homogenate through double layered cheese cloth into a new 50 mL plastic conical tube.
5. Pipette 1.4 mL homogenate into an eppendorf tube (2 mL safe-lock tubes; 02236352, Eppendorf AG, Hamburg, Germany).
6. Centrifuge samples at 1,500 x g for 15 min. at 25C to remove traces of insoluble components.
7. Collect 1mL of supernatent, discard pellet. Place supernatent into 1.5mL eppendorf tube.

Protein Concentration

66. Prepare a concentration series (20-2000 µg/mL) of bovine serum albumin (BSA) using suspension buffer
67. Place 25 µL BSA standards and diluted myofibrillar protein samples on a microwell plate
68. Add 200 µL BCA working reagents (50:1; Reagent A:Reagent B) into respective wells in the microwell plate
69. Mix protein samples and BCA working reagents thoroughly on a plate shaker for 30 sec
70. Incubate the microwell plate at 37°C for 30-40 min and cool to room temperature
71. Read absorbance at **562 nm** on a microplate reader (SpectraMAX 250, Molecular devices, Sunnyvale, California)
72. Protein concentrations are expressed as µg/mL
73. Dilute sample to 6.4mg/mL using whole-muscle solubilizing buffer.
74. Dilute samples to a final protein concentration of 4mg/mL with 50% (volume of original solution in step 8) of gel buffer (3mM EDTA, 3%[wt/vol]SDS, 30%[vol/vol]glycerol, 0.001% pyronin Y [wt/vol], 30mM Tris-HCl, pH 8.0) and 10% of 2-mercaptoethanol.
75. Heat samples for 15 minutes at approximately 50C and stored at -80C.

SDS-PAGE Electrophoresis

Supplies:

15% polyacrylamide separating gels.

Running Buffer (25mM Tris, 192mM glycine, 2mM EDTA, and 0.1%[wt/vol] SDS).

1. Remove the Mini-PROTEAN TGX Precast Gel 4-20% from the package
2. Pull the tape off the bottom, remove the comb
3. Place into electrode assembly with short plate inward
4. Place second gel on the other end of the assembly
5. Lock the two gels in place
6. Place the assembly in the tank
7. Fill the chamber with Bio Rad 1x Tris/Glycine/SDS # 161-0732 running buffer until the short plate is completely covered
8. Add buffer to the outer tank to either 2 or 4 gel line, depending on how many gels you are running
9. Load 10 μ L Bio-Rad Kaleidoscope Pre-stained standards in the first well
10. Load 40ug sample into remaining wells
11. Place lid on tank, aligning color coded banana plugs and jacks
12. Set constant voltage to **360 V**
13. Run until tracking dye in the Laemmle buffer of each sample reaches the black finishing line (30 minutes to an hour)
14. Turn off power supply
15. Remove gel from assembly
16. Remove gel by gently opening key provided in precast gel package

Western Blotting

1. Equilibrate gel in transfer buffer (25 mM Tris-base, 192 mM glycine, 2mM EDTA, and 15% [vol/vol] methanol at pH 9.2) for 20 min
2. Soak precut filter paper (170-3932, Bio Rad Laboratories, Hercules CA) and fiber pads in transfer buffer for 5 min
3. Wet PVDF membranes (IPFL20200, Millipore) in methanol for 15 second and rinse briefly in deionized water before soaking in transfer buffer
4. Prepare the gel sandwich as follows:
 - a. Place the cassette with the black side (anode) down in the transfer buffer in the cassette assembly box
 - b. Place one pre-wetted fiber pad on the black side of the cassette
 - c. Place filter paper on the fiber pad
 - d. Place equilibrated gel on filter paper (roll out all bubbles)
 - e. Place pre-wetted membrane on the gel (roll out all bubbles)
 - f. Place the other filter paper (roll out all the bubbles) and fiber pad respectively
5. Close the cassette firmly without moving the gel and filter paper sandwich and lock the cassette with the white latch
6. Place prepared sets in module
7. Place module in tank and fill with transfer buffer
8. Run Western blot for **90 min** at a constant amperage of **90V**
9. To prevent nonspecific antibody binding, block membranes with PBS-TWEEN [10 mL Odyssey Blocking buffer (927-40100; LI-COR, Lincoln NE) and 0.1% TWEEN 20 (10uL)] containing 5% non-fat dry milk for 60 minutes at room temperature.

Primary Antibody

1. Use Monoclonal Anti-Desmin (D1033: Sigma Aldrich, St. Louis, MO)
Dilute total concentration solution as a 1:40,000 ratio with PBS-TWEEN20 solution.
2. Let rock for an hour at room temperature.
3. Let rock overnight at 4C (~12-18 hours).
4. After primary incubation blots, wash in PBS-Tween 3 time for 10 minute intervals.

Secondary Antibody

1. Reconstitute contents in the original IRDye 680LT conjugated goat anti-mouse IgG1 vial with 0.5 mL sterile distilled water
2. Mix gently by inverting and allow rehydrating for at least 30 min before use (this is stable for three months at 4°C) DO NOT EXPOSE TO LIGHT
 - a. 1 µL in 10 mL Odyssey Blocking Buffer + 0.2% Tween20 (20 µL)
3. Incubate blots in diluted secondary antibody for 60 min at room temperature with gentle shaking (make sure not to expose to light)
4. Pour off secondary antibody solution
5. Wash membrane four times for 10 min each with 15 mL of 1x Tris buffered saline + 0.1% Tween20 on rocking platform
6. Wash membrane with 15 mL of 1x Tris buffered saline for 30 mins to remove residual Tween20

Membrane Imaging

1. Use Odyssey Infrared Imaging System (LI-COR Biosciences, Lincoln NE) at 700 nm channel as integrated intensity
2. Imaging conditions:
 - a. Resolution: 169 nm
 - b. Quality: low
 - c. Focus: offset 0 mm
 - d. Channels: 700 nm
 - e. Intensity: 3.0 or 3.5
 - f. Image size: x-10 y-7

Appendix XV

ISOPROSTANES

Sample Preparation:

1. Immediately after sample collection, powder meat and store in -80°C .
2. After storage, place 30 mg of powdered meat in 5 mL eppendorf tubes.
3. Homogenize sample in 2 mL of 2 N NaOH using Micropolytron. Spin for 20 seconds. (Clean Micropolytron between each sample).
4. Heat homogenized sample in water bath at 45°C for 2 h to ensure hydrolysis.
5. After hydrolysis, cool tissue samples to room temperature (~20 minutes), then neutralize pH using 2 mL of 2N HCl.
6. Split samples into two 2 mL eppendorf tubes. Using microcentrifuge, spin samples at $10,000 \times g$ for 15 min at 4°C .
7. Before Assaying, check to be sure each neutralized sample is in the pH range of **6-8**. If not, adjust the pH to this range by adding 100 μL of the sample to 100 μL of the provided *Neutralization Solution*.
8. Use supernatant for Assay Protocol

Preparation of 8-iso-PGF 2α Standards:

1. Prepare fresh standards by diluting the 8-iso-PGF 2α Standard from 200 $\mu\text{g}/\text{mL}$ to 0.2 $\mu\text{g}/\text{mL}$ in Sample Diluent for a 1:1000 final dilution. (Example: Add 5 μL of 8-iso-PGF 2α Standard stock tube to 4.995 mL of Sample Diluent)
2. Prepare a series of the remaining 8-iso-PGF 2α standards according to Table 1.

Standard Tubes	8-iso-PGF 2α Standard (μL)	Sample Diluent (μL)	8-iso-PGF 2α Standard (pg/mL)
7	1 μL of Standard Stock	4999 μL	40,000
6	250 μL of Tube #1	750 μL	10,000
5	250 μL of Tube #2	750 μL	2,500
4	250 μL of Tube #3	750 μL	625
3	250 μL of Tube #4	750 μL	156.25
2	250 μL of Tube #5	750 μL	39.0625
1	250 μL of Tube #6	750 μL	9.7656
0	0 μL	200 μL	0

Assay Protocol

Note: Each 8-iso-PGF2 α Standard and unknown samples should be assayed in duplicate or triplicate. A freshly prepared standard curve should be used each time the assay is performed.

1a. Immediately before use, dilute the Anti-8-iso-PGF2 α Antibody 1:1000 with Sample Diluent.

1b. Add 100 μ L of the diluted ***Anti-8-iso-PGF2 α Antibody*** to the ***Goat Anti-Rabbit Antibody*** Coated Plate. Incubate 1 hour at 25°C on an orbital shaker.

2. Prepare 300 mL of 1x ***Wash Buffer*** by diluting 10x Wash Buffer Concentrate
- 30 mL 10x Wash Buffer Concentrate
- 270 mL Deionized Water
Warm ***Substrate solution*** to room temperature

3. Make Preparation of Isoprostanes Standards as labeled in **Preparation of 8-iso-PGF2 α** .

4. After incubation, remove the antibody solution from the wells. Wash wells 5 times with 300 μ L 1X Wash Buffer per well. After the last wash, empty the wells and tap microwell plate on absorbent pad or paper towel to remove excess wash solution.

Note: Thorough washing is necessary to remove all of the azide present in the antibody solution.

5a. Immediately before use, dilute the conjugate 1:80 with Sample Diluent. Only prepare enough of the diluted conjugate for the number of wells immediately used.

5b. Combine 55 μ L of the ***8-iso-PGF2 α standard*** or sample and 55 μ L of ***8-iso-PGF2 α -HRP conjugate*** in a microtube and mix thoroughly. Transfer 100 μ L of the combined solution per well. A well containing Sample Diluent can be used as a control. Incubate 1 hour at 25°C on an orbital shaker.

6. After incubation, remove the combined solution from the wells. Wash 5 times with 300 μL of 1X Wash Buffer per well. After the last wash, empty wells and tap microwell plate on absorbent pad or paper towel to remove excess wash solution.
7. Add 100 μL of ***Substrate Solution*** to each well. Incubate at room temperature for 15 minutes on an orbital shaker.
8. Stop the enzyme reaction by adding 100 μL of ***Stop Solution*** to each well. Results should be read immediately (color will fade over time).
9. Read absorbance of each well on a microplate reader using 450 nm as the wavelength.